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EUROPEAN PATENT APPLICATION

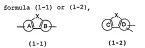
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- (54) Conjugated polymer comprising dibenzothiophene- or dibenzofuran-units and their use in polymer LEDs
- (57) Provided is a polymer compound having a polystyrene reduced number average molecular weight of 103-108, and comprising a repeating unit of formula (1-1) or (1-2):



wherein Ring A, Ring B, Ring C, and Ring D each independently represent an aromatic ring, and X represents S or O. The compound is usable as a light-emitting material, a charge transporting material, etc.

Description

BACKGROUND OF THE INVENTION

5 Field of the Invention

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[0001] The present invention relates to a polymer compound and a process for producing the same. Specifically, the invention relates to a polymer compound which is expectable as a material for a polymer light emitting device (hereinafter may be referred to as polymer LED), and a process for producing the same.

Description of the Related Art

[0002] Unlike low molecular weight it, a high molecular weight light-emitting material ferein, light-emitting material presents are material having fluorescence and phosphoroscence) and a high molecular weight charge transporting material are soluble in a solvent, and are variously studied from the ability of the organic layer in a light-emitting device to be formed by the application method. As a polymer compound which can be used for electronic devices, such as polymer LED, as a light-emitting material or a charge transporting material, polyphenylene vinylene derivatives, poly fluorene derivatives, polythorous derivatives, contributions ordivatives, overhickness, and as a contribution of the device of the derivative solvent or the contribution of the device of

[0003] The object of the present invention is to provide a new polymer compound usable for a light-emitting material, or a charge transporting material, etc., a process for producing thereof, and a polymer light-emitting device using said polymer compound.

SUMMARY OF THE INVENTION

[0004] As a result of extensive studies in order to solve the above problems, the present inventors found that a polymer compound having a polystyrene reduced number average molecular weight of 10³-10⁸, and comprising a repeating unit represented by the below formula (1-1) or (1-2).

is usable as a light-emitting material, a charge transporting material, etc. and completed the present invention.

[0005] The polymer compound of the present invention comprises one or more kinds of repeating unit represented by the above formula (1-1) or (1-2).

[0006] In the above formula (1-1) or (1-2), Ring A, Ring B, Ring C, or Ring D each independently represent an aromatic ring. X represents S or O.

DETAILED DESCRIPTION OF THE INVENTION

[0007] As the aromatic ring, exemplified are: aromatic hydrocarbon rings such as a benzene ring, naphthalene ring, anthracene ring, tetracener ring, pentacener ring, repron ring, and phenanthrener ring, etc; and heterocyclic aromatic rings such as a pyridine ring, blyriddine ring, phenanthreoline, quinoline ring, isoquinoline ring, thron ring, pyrrole infing, dec.

[0008] The repeating unit containing structure represented by the above formula (1-1) or (1-2), may have a substituent selected from the group, consisting of an alkyl group, alkoys group, alkylling group, anylong group, and group, anylong group, anylong group, and group, and group, and group, and group, and group, and group, anylong group, and group, an

[0009] Here, the alkly group may be any of linear, branched or cyclic, and usually has about 1 to 20 carbon atoms, preferably 3 to 20, and specific examples thereof include methyl group, lothyl group, propyl group, i-propyl group, butyl, I-butyl, I-butyl, pennyl group, heavyl group, cyclohexyl group, heptyl group, podyl group, 2-ethylhexyl group, nonyl group, decyl group, 3,7-dimethyloctyl group, lauryl group, trifluoromethyl group, pentalluoroethyl group, pentluorobutyl, pentfluorothexyl group, pentluoroethyl group, eth. Pentluorothexyl group, cohyl group, cell-withpoxyl group, decyl group, and cell group.

- 3,7-dimethyloctyl group is preferable.
- [0010] The alkoxy group may be any of linear, branched or cyclic, and usually has about 1 to 20 cation atoms, preferable 3 to 20.5 Apedific assurghes thereof include methoxy group, behoty group, prolypoxy group, butory group, i-butoxy group, i-butoxy group, pentyloxy group, heayloxy group, cycloheayloxy group, heptyloxy group, pentyloxy grou
- [0011] The alkythio group may be any of linear, branched or cyclic, and usually has about 1 to 20 carbon atoms, preferably 3 to 20. Specific examples thereof include methythio group, athythio group, preythio group, preythio group, braythio group, preythio group, at-yithio group, cctylthio group, 2-ethythreythio group, octylthio group, at-yithio gro
- 15 [0012] The anyl group usually has about to 60 carbon atoms, preferably 7 to 48. Specific examples thereof include a phenyl group, C-φ₂₂ allowphenyl group, C-φ₂₂ aneans the number of carbon atoms is 1o 12; Q-G₁₂ allyphenyl group, 1-naphthyl group, 2-naphthyl group, 1-anthracenyl group, 2-anthracenyl group, 9-anthracenyl group, pentalfluorophenyl group, etc. Q-G₁₂ allowphenyl group, and G-G₁₂ alloyhpenyl group are preferable. The anyl group is atomic group in which a hydrogen atom is removed from an aromatic hydrocarbon. The aromatic hydrocarbon includes the properties of the properti
 - [0013] Specifically, examples of C₁-C₁₂ alkoxyphenyl include methoxyphenyl, ethoxyphenyl, propyloxyphenyl, i-propyloxyphenyl, butoxyphenyl, i-butoxyphenyl, i-butoxyphenyl, pentyloxyphenyl, hasyloxyphenyl, cyclohoxyl oxyphenyl, heptyloxyphenyl, octyloxyphenyl, 2-ethylhexyloxyphenyl, nonyloxyphenyl, decyloxy, 3,7-dimethyloctyloxyphenyl, etc.
 - [0014] Examples of CrC12 alkytyhenyl group include a methylphenyl group, ethylphenyl group, dimethylphenyl group, propylphenyl group, mestyl group, methylethylphenyl group, propylphenyl group, butylphenyl group, pentylphenyl group, pentylpheny
- 30 [0015] The aryloxy group usually has about 6 to 60 carbon atoms, preferably 7 to 48. Specific examples thereof include a phenoxy group, C-p_{C-R} alkoxyphenoxy group, p-C-p_{C-R} alkoyphenoxy group, and C-p_{C-R} alkoyphenoxy group, and and and alkoyphenoxy group, and and alkoyphenoxy group, and and alkoyphenoxy group, alkoyphenoxy group, and alkoyphenoxy group, and alkoyphenoxy group, alkoyphenoxy group, alkoyphenoxy group, and alkoyphenoxy group, alkoyphenoxy group, alkoyphenoxy group, alkoyphenoxy group, alkoyphenoxy group, alkoyphenoxy group, alko
 - [0016] Specific examples of C₁-C₁₂ alkylphenoxy group include a methylphenoxy group, ethylphenoxy group, Jimethylphenoxy group, propylphenoxy group, 13,5-trimethylphenoxy group, methylethylphenoxy group, i-burlylphenoxy group, i-burlylphenoxy group, propylphenoxy group, i-burlylphenoxy grou
- phenoxy group, etc.

 [0017] The anythol group usually has about 6 to 60 carbon atoms, preferably 7 to 48. Specific examples thereof include a phenythic group, C₁-C₁₂ alkoxyphenythic group, C₂-C₁₂ alkoxyphenythic group, C₃-C₁₄ alkoxyphenythic group, and C₄-C₁₂ alkoy phenythic group, and C₄-C₁₂ alky phenythic group, and C₄-C₁₂ alky phenythic group are preferable.
 - [0018] The anylalkyl group usually has about 7 to 60 carbon atoms, preferably 7 to 48. Specific examples thereof include a phenyi-C₁-C₂ alkyl group, C₁-C₂ alkyl group, C₁-C₂ alkyl group, 2-naphtyl-C₁-C₁₂ alkyl group are preferable.
- [0019] The anyl alkoy group usually has about 7 to 60 carbon atoms, preferably 7 to 48. Specific examples thereof include phenyl-C₁-C₁₂ alkoy groups such as a phenylmethoxy group, phenyleuthoxy group, phenylmethoxy group, clac.; C₁-C₁₂ alkoy group, C-1-C₁₂ alkoy group, and C-1-C₁₂ alkoy group are preference on C
 - [0020] The aryl alkylthio group usually has about 7 to 60 carbon atoms, preferably 7 to 48. Specific examples thereof include a phenyl-C₁-C₁₂ alkylthio group, C₁-C₁₂ alkylthio group alkylthio group, C₁-C₁₂ alkylthio group alkylthi

alkylthio group, 1-naphtyl- C_1 - C_{12} alkylthio group, 2-naphtyl- C_1 - C_{12} alkylthio group, etc. C_1 - C_{12} alkoxyphenyl- C_1 - C_{12} alkylthio group, and C_1 - C_{12} alkylphenyl- C_1 - C_1 2 alkylthio group are preferable.

[0021] The aryl alkenyl group usually has about 7 to 60 carbon atoms, preferably 7 to 48. Specific examples thereof include a phenyl-C₂-C₁₂ alkenyl group, C₁-C₁₂ alkenyl group, C₁-C₁₂ alkenyl group, C₁-C₁₂ alkenyl group, 2-naphlyl-C₂-C₁₂ alkenyl group, etc. C₁-C₁₂ alkenyl group, and C₂-C₁₂ alkenyl group are preferable.

[0022] The any falkymy group usually has about 7 to 80 carbon atoms, preferably 7 to 48. Specific examples thereof include a pheny-le₂-Ct₁₂ alkymyl group, Ct₁-Ct₁₂ alkymyl group, Ct₁-Ct₂ alkymyl group, Ct₁-Ct₂ alkymyl group, 1-maphhyl-Ct₂-Ct₁₂ alkymyl group, 1-maphhyl-Ct₂-Ct₂-Maphhyl-Ct₂

and G₁-C₁₂ alkjohenyl-C₂-C₁₂ alkynyl group are preferable.

[0023] The substituted amino group means an amino group which has one or two substituents selected from an alkyl group, anylaikyl group, or monovalent heterocyclic group. The alkyl group, anylaikyl group, or monovalent heterocyclic group. The alkyl group, anylaikyl group, or monovalent heterocyclic group. The alkyl group, anylaikyl group, or monovalent heterocyclic group. The alkyl group, anylaikyl group, or monovalent heterocyclic group. The alkyl group, anylaikyl group or monovalent heterocyclic group.

monovalent heterocyclic group may have a substituent. The number of carbon atoms is usually about 1 to 60 without including the carbon atoms of the substituent, and preferably 2 to 48.

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5 [0024] Specifically exemplified are: a methylamino group, dimethyl amino group, ethylamino group, fiethylamino group, bropylamino group, cyclohexylamino group, pethylamino group, cyclohexylamino group, pethylamino group, grou

[0028] The substituted silyl group means a silyl group which has 1 to 3 substituents selected from an alkyl group, anyl group, aryl group, arrylalkyl group, or monovalent heterocyclic group. The number of carbon atoms is usually about 1 to 80, and preferably 3 to 48. The alkyl group, arylalkyl group, or monovalent heterocyclic group may have a

substituent. [0026] Specifically exemplified are: a trimethylally group, triethylally group, tripropylally group, tri-propylally group, triethylally group, tripropylally group, triethylally group, triethylally group, tripropylally group, triethylally group, g

[0027] As the halogen atom, a fluorine atom, chlorine atom, bromine atom, and an iodine atom are exemplified.
[0028] The acyl group usually has about 2 to 20 carbon atoms, preferably 2 to 18. Specific examples thereof include an acetyl group, propionly group, butrynl group, isobutrynl group, pivaloyl group, benzoyl, trifluoroacetyl group, pentalluorobenzoyl, etc.

[0029] The acyloxy group usually has about 2 to 20 carbon atoms, preferably 2 to 18. Specific examples thereof include an acetoxy group, propion/toxy group, butyloxy group, isobutyyloxy group, pivaloyloxy group, benzeyloxy group, frill/proacetyloxy group, pentafluorobenzoyloxy group, etc.

[0030] The imino group has about 2 to 20 carbon atoms, preferably 2 to 18. Specific examples thereof include the groups represented by the following structural formulae, and the like.

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[0031] The amid group usually has about 21 o 20 carbon atoms, preferably 2 to 18. Specific examples thereof include formamide group, eatennide group, proploamied group, butyor amide group, butwo amide group, filteronaceatemide group, pentafluorobenzamide group, discontamide group, disc

25 [0032] The imide group usually has about 2 to 60 carbon atoms, preferably 2 to 48. Specific examples thereof include the groups shown below.

[0033] A monovalent heterocyclic group means an atomic group in which a hydrogen atom is removed from a heterocyclic compound, and usually has about 4 to 80 carbon atoms, preferably 4 to 20. The carbon atoms of the substituent are not counted as the number of carbon atoms of the heterocyclic group. The heterocyclic compound means an organic compound having a cyclic structure in which at least one heteroatom such as oxygen, sulfur, nitrogen, phosphorus, brone, faci. s contained in the cyclic structure as the element other than carbon atoms.

[0034] Specific examples thereof include a thienyl group, C_1 - C_{12} alkylithienyl group, pyrotyl group, furyl group, pyridyl group, C_1 - C_1 - C_1 - C_2 - C_1 - C_1 - C_2 - C_1 - C_1 - C_2 - C_2 - C_3 - C_3 - C_4 -

[035]. The substituted carboxyl group usually has about 2 to 60 carbox atoms, preferably 2 to 46. The substituted carboxyl group means a carboxyl group having a substitutent such as an alkyl group, anyl group, anyllally group, or monovalent heterocyclic group. Specific examples thereof include a methoxycarbonyl group, ethoxycarbonyl group, public propopycarbonyl group, propopycarbonyl gr

[0036] In the above examples of substituents, the substituent containing an alkyl chain may be linear, branched or cyclic one, or the combalion thereof. A risk alkyl chain which is not linear, exemplified are isoamy group, 2-dityhbasyl group, 8-dimethylocyl group, exclobely group, 4-diversely group, excl. Moreover, two alkyl chain ends may be connected to form a ring. Furthermore, methyl or ethyl as a part of said alkyl chain may be replaced by a group containing a hetero acm, or a methyl or ethyl group which is substitued with one or more fluorine atoms. Here, as the hetero atom, an away and a sufficient or a sufficient production of the sufficient production and the sufficient production and a sufficient production and

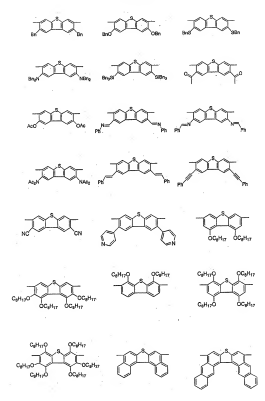
[0037] Among the structures represented by formula (1-1), concrete examples in case of X=S include the followings.

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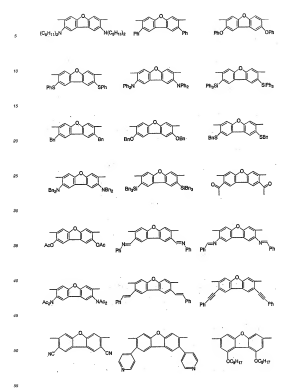
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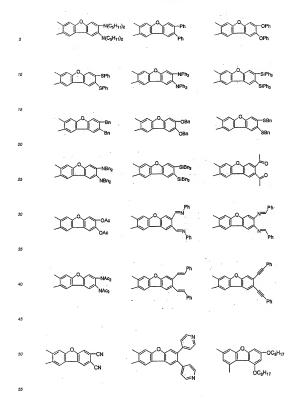
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[0038] Among the structures represented by formula (1-1), concrete examples in case of X=O include the followings.

ε



5	N(C ₆ H ₁₁) ₂ N(C ₆ H ₁₁) ₂	SPh	S OPh OPh
10	SPh	NPh ₂	SiPh ₃
15	181		
20 _	S Bn Bn	SOBN	SBn
25	NBn ₂	SiBn ₃	TS-C-0
30		P h	Ph
35	S OAc	S N	S N Ph
40	\	Ph	Ph
45	NAC ₂	Ph	Ph
50	\	s . D	
55	CN	III	OC ₈ H ₁₇



[0041] In the formulae, Me represents a methyl group, Ph represents a phenyl group, Bn represents a benzyl group, and Ac represents an acetyl group.

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[0043] Among them, an alkyl group, alkovy group, alkylthio group, aryl group, aryloxy group, arylishlo group, arylakylthio group, substituted alkyl group, substituted skyl group, fundrie atom, avgl group, arylakynthio group, midde group, arylakyl group, askovy group, arylakyl group, arylakkyl group, arylakyl group,

[0044] As the repeating unit in the polymer compound of the present invention, a repeating unit represented by formula (1-1) is preferable.

[0045] As for Ring A, Ring B, Ring C, and Ring D, those containing only an aromatic hydrocarbon ring are preferable. [0046] Among them, preferable are those which have a structure represented by the above formula (1-1), and have round represented by the above formula (1-1), and have represented by the above formula (1-1), and have represented by the above formula (1-1), and have represented by the representation of the preferable are the repeating units selected (from the below formulas (2-1), (2-2), (2-3), (2-3), (2-4), and (2-5).

$$(R_1)_a$$
 $(R_2)_b$

(Ra)c (Ra)a

(2-2)

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(R₆)_e (R₆)_t

(2-3)

(F₇)_g (F₀)_h

(2-4)

(2-5)

[0047] In the formulae, X represents S or O. R., R.g., R.g.,

[0049] In the above formulae (2-4) and (2-5), the substituents R₇ to R₁₀ may substitute on any carbon atoms of two benzene rings constituting the condensed aromatic ring.

[0050] In view of improving the solubility of polymer compound, changing the wavelength of light-emission, and improving the efficiency of light-emission, it is preferable that a+b of formula (2-1), o+d of formula (2-2), e+f of formula (2-3), e+f of formula (2-4), e+f of formula (2-

[0051] When R_1 exists in plural, a plurality of them may be the same or different. As for R_2 , R_3 , R_4 , R_5 , R_6 , R_7 , R_8 , and $R_{\rm in}$, when they exist in plural, respectively, a plurality of them may be the same or different.

[0052] When X-S, among the repeating units selected from the above (2-1), (2-2), (2-3), (2-4) and (2-5), the repeating unit represented by formula (2-1) is preferable but all ray and/or P₀ is an alityl group having 3 or more carbon atoms, an alloxy group having 3 or more carbon atoms, an alloxy group having 3 or more carbon atoms, an alloxy group, anyloxy gro

monovalent heterocyclic group.

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[0053] When X=O, among the repeating units selected from the above (2-1), (2-2), (2-3), (2-4) and (2-5), the repeating unit represented by formula (2-6) is preferable.

In the formula, R_1 and R_2 each independently represent an alkyl group having 3 or more carbon atoms, alkoxy groups having 3 or more carbon atoms, alkylthio groups having 3 or more carbon atoms, any group, anyloxy group, anylakyl group, anyloxy group, anyloxy group, parylakivly group, anyloxy group, parylakivly group, anyloxy group, parylakivly gr

[0054] Total amount of the repeating units represented by the above formula (1-1) and (1-2), is usually 1% by mole or more and 100% by mole or more, and more preferably 90% by mole or more, and more preferably 90% by mole or more and 90% by mole or flow. In the preferably 10% by mole or more and 100% by mole or less. When X-0, it is preferably 10% by mole or more and 100% by mole or less, and more preferably 50% by mole or some and 90% by mole or less.

[0055] In view of improving fluorescence intensity, when a copolymer consisting of repeating units represented by either one of formula (1-1) or formula (1-2), and the repeating units have only the structure of either X=5 or X=O, it is preferable that the repeating units are two or more monomers sharing different fixed of substitutents.

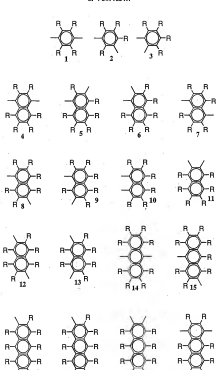
[0056] The polymer compound of the present invention is preferably a copolymer which contains the repeating unit having structure of the above formula (1-1) and/or (1-2) and one or more of the other repeating unit.

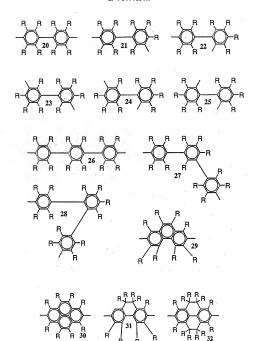
[0057] Examples of the other repeating unit than those represented by formulas (1-1) and (1-2) preferably include the repeating units represented by the below formula (3), formula (4), formula (5), or formula (6).

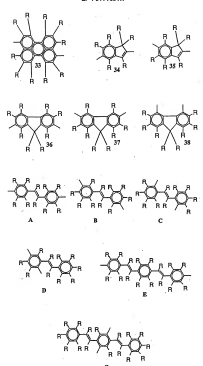
[0058] In the formulae, A_{11} , A_{12} , A_{13} , and A_{14} represent each independently an arylene group, a divalent heterocyclic group, or a divalent group having metal-complex structure. X_1 , X_2 , and X_3 represent each independently $-CH_{11}-CH_{12}-C$

[0059] In the present invention, the arylene group is an atomic group in which two hydrogen atoms are removed from an aromatic hydrocarbon, and usually has 6 to 60 carbon atoms, preferably 6 to 20 carbon atoms. The aromatic hydrocarbon also includes those containing a condensed ring, and two or more of independent benzene rings or condensed rings bonded through a group such as a direct bond, a vinylene group or the like.

[0060] As the arytene group, exemplified are phenylene group (for example, formulas 1-3), naphthalenediyl group (following formulas 1-14), phipharyl-dyl group (following formulas 1-14), phipharyl-dyl group (following formulas 20-25), fluorene-dyl group (following formulas 36-38), terpharyl-dyl group (following formulas 26-28), stilbene-dyl group (following formulas 26-28), stilbene-dyl group (following formulas 28-38) etc. Among them, phenylene group, biphenylene group, fluorene-dyl group, and stilbene-dyl group are perforable.







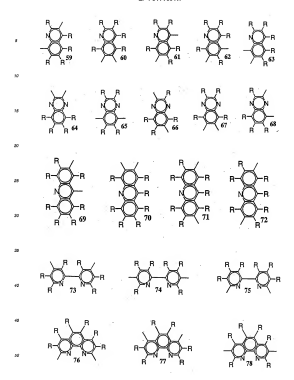
5 [0061] In the present invention, the divalent heterocyclic group means an atomic group in which two hydrogen atoms are removed from a heterocyclic compound, and the number of carbon atoms is usually about 3 to 60.
[0062] The heterocyclic compound means an organic compound having a cyclic structure in which at least one heteroatom such as oxygen, sulfur, nitrogen, phosphorus, boron, arsenic, etc. is contained in the cyclic structure as the

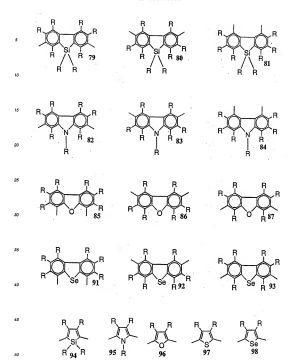
element other than carbon atoms.

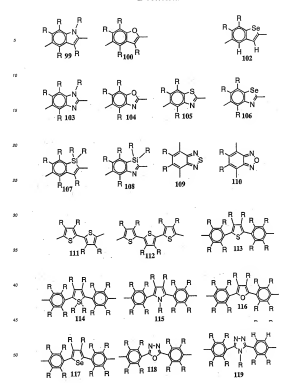
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[0063] As the divalent heterocyclic compound group, followings are exemplified.

[0064] Groups containing a nitrogen as a hetero atom; pyrkdime-dyl group (following formulas 39-44), diazaphonylene group (following formulas 45-48), quinolinedyl group (following formulas 45-48), authority formulas 45-48), quinolinedyl group (following formulas 45-48), authority formulas 45-48), principal group (following formulas 73-75), phenarchicoline dyl group (following formulas 73-75), phenarchicolined group containing a hetero atom, such as silicon, nitrogen, suffur, selenium, etc. (following formulas 79-39); formerbered-ring heterocyclic compound groups containing a hetero atom such as silicon, nitrogen, suffur, selenium, etc. (following formulas 94-98); formerbered-ring heterocyclic compound groups containing a hetero atom such as silicon, nitrogen, suffur, selenium, etc. (following formulas 90-108); groups in which 5 membered ring heterocyclic compound group containing silicon, nitrogen, suffur, selenium, etc. as a hetero atom is connected with a phenyl group at the a position of the hetero atom is connected with a phenyl group at the a position of the hetero atom is connected with a phenyl group at the a position of the hetero atom is connected with a phenyl group at the a position of the hetero atom is connected with a phenyl group at the a position of the hetero atom is connected with a phenyl group at the a position of the hetero atom is connected with a furyl group, or thienyl group (following formulas 113-119); and groups in which 5 membered ring heterocyclic compound group containing oxygen, nitrogen, sulfur, etc. as a hetero atom is connected with a furyl group, or thienyl group (following formulas 110-15).







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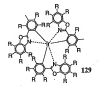
[0065] In the present invention, the divalent group having metal-complex structure is a divalent group in which two hydrogen atoms are removed from the organic ligand of the metal complex.

[0068] The number of carbon atoms of the organic ligand is usually about 4 to 60. Examples of the organic ligand include 9-quincinol and its derivatives, percepul-organic and its derivatives, 2-phen-yp-included and its derivatives, 2-phen-yp-included and its derivatives, 2-phen-yp-included and its derivatives, 2-phen-yp-included and its derivative, etc. are exemplified.

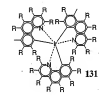
[0067] As the central metal of the complex, for example, aluminum, zinc, beryllium, iridium, platinum, gold, europium, terbium, etc. are exemplified.

[0068] As the metal complex having an organic ligand, exemplified are a metal complex and a triplet luminescence complex which are known as a low molecular weight fluorescent material or a phosphorescent material.

[0069] Specifically as the divalent group having metal-complex structure, the followings (126-132) are exemplified.









[0070] In the example shown by the above formulas 1-132, Represents each independently a hydrogen atom, allyly group, alkoxy group, allylythio group, anylathyn group, substituted arisen group, anylathyn group,

[0071] The carbon atom contained in the group represented by formulas 1-132 may be replaced with a nitrogen atom, oxygen atom, or sulfur atom, and the hydrogen atom may be replaced with a fluorine atom.

[0072] Among the repeating units represented by formula (3), the repeating units represented by below formula (7), (8), (9), (10), (11), or (12) are preferable.

In the formula, R₁₈ represents an alkyl group, alkoxy group, alkythio group, any group, anyloxy group, anylish group, anylak y group, anyloxy group, animo group, sibstituted amino group, sily group, pathol group, animo group, sily group, animo g

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$$\begin{pmatrix} R_{17} \\ R_{17} \\ R_{18} \end{pmatrix}_{n}$$
(8)

in the formula, R₁₇ and R₁₈ represent each independently an alkyl group, alkoxy group, alkylthio group, anyla group, anino group, anino group, anino group, anino group, anide group, imino group, anono group, anide group, imino group, anide group, imino group, anide group, anonovalent heterocyclic group, carboxyl group, substituted carboxyl group, or cyano group, mand n represent each independently an integer of 0-3. When R17 and R18 exist in plural, respectively, a plurally of them may be the seame or different.

In the formula, R₁₉ and R₂₂ represent each independently an alkyl group, alkoxy group, alkylthio group, aryl group,

anyloxy group, anylativg group, anylati

$$-\left(A_{f_{13}}\right)_{X}^{X_{4}} \left(A_{f_{14}}\right)_{y} \left(A_{g_{23}}\right)_{q}$$

$$\left(B_{23}\right)_{q}$$

$$\left(10\right)$$

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In the formula, R_{23} represents an alkyl group, alkoxy group, alkythio group, aryl group, aryl group, arylathog group, arylathog group, arylathyl group, a

X4 represents O, S, SO, SO2, Se, or Te. When R₂₃ exists in plural, a plurality of them may be the same or different.

In the formula B_{Ba} and B_{Ba} each independently represent an alkyl group, alkxoy group, alkythio group, and y group, and group, and y group, and y

$$\begin{array}{c}
\begin{pmatrix}
R_{30} \\
T
\end{pmatrix}_{t} \\
R_{31} \\
R_{32} \\
R_{34} \\
\begin{pmatrix}
R_{36} \\
R_{36}
\end{pmatrix}_{u}$$
(12)

In the formula, R₃₀ and R₃₅ each independently represent an alkyl group, alkoxy group, alkylthio group, aryl group,

aryloxy group, arylthio group, arylalkyl group, arylalkyny group, arylalkynthio group, arylalkynyl group, ar

[0073] Among the repeating units represented by the above formula (4), the repeating unit represented by the below formula (13) is preferable.

In the formula, $A_{\rm fb}$, $a_{\rm fb}$,

(13)

[0074] In the above formula, R is the same as those of the above formulas 1-132. In formulas 1-140, a plurality of Rs are contained in one structural formula, and they may be the same or different. In order to improve the solubility in a solvent, it is perferable to have one or more substituents other than hydrogen, and it is more preferable that the symmetry of the repeating unit including the substituent is little. Furthermore, when R contains an anyl group or a heterocyclic group as a part of the above formula. It may further have one more or more substituents.

[0075] 'In the above examples of substituent Roontaining an alkyl chain as a part thereof, they may be linear, branched or cyclic one, or the combination thereof. As the alkyl chain which is not linear, exemplified are isoamyl group, 2-ethyl-hexyl group, 3,7-dimethylocyl group, cyclohexyl group, 4-Cy-Cy₂ alkylcyclohexyl group, etc. In order to improve the solubility of the polymer compound into a solvent, it is preferable that a linear or branched alkyl chain is contained in one or more substituents.

[0076] Moreover, a plurality of Rs may be connected to form a ring. Furthermore, when R contains an alkyl chain, said alkyl chain have be interrupted by a group containing a hetero atom. Here, as the hetero atom, an oxygen atom, a sulfur atom, a nitrogen atom, etc. are exemplified.

[0077] Among them, the repeating units represented by the below formula (13-2) is preferable.

In the formula, R₅₀, R₅₁, and R₅₂ represent each independently an alkyl group, alkoxy group, alkythio group, arylakryngroup, hi, ii, and ji each independently represent an integer of 0-4. 2 represents an integer of 0-2.
When R₅₀, R₅₁, and R₅₂ exist in plural, respectively, a plurally of them may be the same or different.

[0078] The polymer compound of the present invention may contain a repeating unit other than the repeating unit represented by the above formula (1-1), (1-2) and formulas (3)-(13) within a range of not injuring light-inversion characteristics or charye transportation characteristics. Moreover, these repeating units and other repeating units may be connected with non-conjugated unit, and the non-conjugated portions thereof may be contained in the repeating units. As the connected structure, the followings and a combination of two or more of them are exemplified. Here, R is a group selected from the same substituents as the above-mentioned, and Ar represents a hydrocarbon group of 6-80 carbon alons.

5 [0073] The polymer compound may also be a random, block or graft copolymer, or a polymer having an intermediate structure thereof, for example, a random copolymer having block properly. From the viveypint for obtaining a polymer compound having high fluorescent or phosphorescent quantum yield, random copolymers having block properly and block or graft copolymers are more preferable than complete random copolymers. Further, the polymer compound may set the property of the polymer compound may be preferable than complete random copolymers. Further, the polymer compound may set the property of the

have a branched main chain and more than three terminals. Moreover, the polymer compound may be a dendrimer.
[0080] Furthermore, the end group of polymer compound may also be protected with a stable group since if a present primerization active group remains intact, there is a possibility of reduction in light entitling property and life-time when made into an device. Those having a conjugated bond continuing to a conjugated structure of the main chain are preferable, and there are exemplified structures connected to an anyli group or heterocyclic compound group via a carbon-carbon bond. Specifically, substituents of the chemical formula 10 in JP 9-45478A are exemplified.

[0081] The polystyrene reduced number average molecular weight of the polymer compound of the present invention is 10³-10⁸, and preferably 10⁴-10⁶.

[0082] As a good solvent for the polymer compound of the present invention, exemptified are chloroform, methylene chloride, dichloroethane, tetrahydrofuran, foluene, xylene, mesitylene, tetrafin, decalin, n-butylbenzene, etc. It can be dissolved in the solvent usually 0.1% by weight or more, although it depends on the structure and the molecular weight of the polymer compound.

[0083] Next, the process for producing the polymer compound of the present invention is explained. The polymer compound of the present invention can be produced by the below method.

56 (Method A) Carrying out condensation polymerization of a compound having two substituents capable of condensation polymerization, corresponding formula (1-1) or (1-2). In case of X-5, the polymer compound can be produced by: produced by: (Method S) Carrying out condensation polymerization of a compound having two substituents whose X-SO_capable of condensation polymerization, corresponding formula (1-1) or (1-2), and then reducing the obtained polymer; or (Method C) Carrying out condensation polymerization of a compound having two substituents whose X-SO capable of condensation polymerization, corresponding formula (1-1) or (1-2), and then reduction the obtained polymer.

[0084] For example, the polymer compound of the present invention can be produced by carrying out condensation polymerization with using a compound represented by Formula (14) as one of raw materials,

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wherein, U represents a repeating unit represented by the above formula (1-1) or (1-2), and Y₁ and Y₂ each independently represent a substituent capable of condensation polymerization. (Method A)

[0085] By using compounds represented by the below formula (15-1), (15-2), and (15-3), as a compound of the above formula (14), polymer compounds having a repeating unit represented by the below formula (15-4), (15-5) and (15-6), and be obtained; respectively.

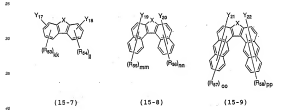
(wherein, $Y_{\rm th}, Y_{\rm th}, Y_{\rm th}, Y_{\rm th}, and <math>Y_{\rm th}$ each independently represent a substituent capable of condensation polymerization. X openeants S or S $R_{\rm 30}$, $R_{\rm 30}$, $R_{\rm 30}$, and $R_{\rm 47}$, represent each independently an eldy group, a naticoxy group, allydithio group, any group, and group, group, and group, group, and group, and group, group, and group, and group, g

[0086] In the above formula (15-2), the substituents Y₁₃, Y₁₄, R₃₈ and R₈₉ may substitute on any carbon atoms of two benzene rings constituting the condensed aromatic ring.

[0087] In the above formula (15-3), the substituents Y₁₅, Y₁₆, R₄₀ and R₄₁ may substitute on any carbon atoms of three benzene rings constituting the condensed aromatic ring.

[0088] Examples of the compounds represented by the above formula (15-1), (15-2), or (15-3) include compounds represented by below formula (15-7), (15-8) or (15-9),

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wherein, Y₁₇, Y₁₈, Y₃₉, Y₂₄, and Y₂₂ each independently represent a halogen atom, allof sulfonate group, any sulfonate group, any sulfonate group, any sulfonate group, any sulfonation methyl group, bnosphonium methyl group, bnosphonium methyl group, bnosphonium methyl group, and phosphonium methyl group, and phosphonium for group, and year group, any group, any group, any group, any group, any sulfonation group, and sulfonation group, any sulfonation group, and sulfon

[0089] In the above formula (15-8), the substituents Y₁₆, Y₂₀, R₅₅ and R₅₆ may substitute on any carbon atoms of two benzene rings constituting the condensed aromatic ring.

[0090] In the above formula (15-3), the substituents Y₂₁, Y₂₂, R₅₇ and R₅₈ may substitute on any carbon atoms of three benzene rings constituting the condensed aromatic ring.

[0091] In the case of X=S, a compound represented by the below formula (15-10) is further preferable.

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wherein, V_{20} and V_{24} represent a bromine atom, chlorine atom, akly sulfonate group, any sulfonate group, any sulfonate group, any sulfonate group, both cester group, sulfonatine methy group, phosphonate methyl group, and boric acid group. Reg. and Ry₂₀ each independently represent an alkyl group having 3 or more carbon atoms, an alkylding group having 3 or more carbon atoms, and skylding group having 3 or more carbon atoms, and skylding group having 3 or more carbon atoms, and group, anylokylding group, anylo

Moreover, in case of X=O, a compound represented by the below formula (15-11) is further preferable.

wherein, Y₂₅ and Y₂₆ represent a bromine atom, chlorine atom, alkyl sulfonate group, aryl sulfonate group, arylaikyl sulfonate group, bofe ester group, sulfonium methyl group, phosphonium methyl group, phosphonium methyl group, and borie cadie group. B-y and B-y, each independently represent an alkyl group having 3 or more carbon atoms, an alkoxy group having 3 or more carbon atoms, an alkoxy group having 3 or more carbon atoms, an alkylthio group having 3 or more carbon atoms, anyl group, arylaikyl group, arylaikyl group, arylaikylthio group, substituted amino group having 4 or more carbon atoms, and substituted alth group having 4 or more carbon atoms.

[0092] Moreover, in case of X=S, the polymer compound of the present invention can be produced by carrying out condensation polymerization of a compound represented by the below formula (22), and then reduction thereof. (Method B)

40 wherein, V represents a repeating unit having a structure of X=SO corresponding to the above U. Y₁ and Y₂ are the as above-mentioned.

[0093] The reduction can be performed according to the below mentioned method of the producing process of compound (14) from a compound of formula (22). The repeating unit V having thiophene-sulfone structure corresponding to each U is as follows.

[0094] For example, by carrying out condensation polymerization of a compound of below formula (15-12) as a compound of formula (22-1), then reducing the product with using a reducing agent, a polymer compound having a repeating unit represented by the above formula (15-1) can be obtained.

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wherein, Y₁₁, Y₁₂, R₃₈, R₃₉, aa, and bb are the same as above-mentioned. [0095] The polymer compound of the present invention can be obtained also by carrying out condensation polymerization of a compound of formula (23), and then reducing the product. (Method c)

Formula (23)

Y1-W-Y2

wherein, W represents a repeating unit having a structure represented by X=SO corresponding to the above U. Y_1 and Y_2 are the same as above-mentioned.

[0096] The repeating-unit W having thiophene structure corresponding to each U is as follows.

[0097] For example, by carrying out condensation polymerization of a compound of below formula (15-13) as a compound of formula (26-1), then reducing the product with using a reducing agent, a polymer compound having a repeating unit preprisent on the provision of the production of the provision of the provisio

wherein, Y11, Y12, R38, R39, aa, and bb are the same as above-mentioned.

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[0098] In the process of the present invention, as the substituent capable of condensation polymerization, exemplified are a halogen atom, affyl sulfornate group, anysitionate group, anysitiony, border ster group, but on the group, process group, process group, process group, process group, process group, bendering process, under the group, process group, process group, proving process, process group, proving proup, proving proving

[0099] Here, as the alkylsulfonate group, exemplified are methane sulfonate group, ethane sulfonate group, trifluoromethane sulfonate group, etc. As the anylsulfonate group, exemplified are benzene sulfonate group, p-toluene sulfonate group, etc. As the anylsulfonate group, exemplified are benzylsulfonate group etc.

[0100] As the boric ester group, exemplified are the groups represented by the below formula.

In the formula, Me represents methyl group and Et represents ethyl group.

[10101] As the sulfonium methyl group, exemplified are the groups represented by the below formula.

(X represents a halogen atom and Ph represents phenyl group.)

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[0102] As the phosphonium methyl group, exemplified are the groups represented by the below formula.

[0103] As the phosphonate methyl group, exemplified are the groups represented by the below formula.

-CH₂PO₂(OR') (X represents a halogen atom and R' represents an alkyl group, aryl group, or arylalkyl group.)

5 [0104] As the monohalogenated-methyl group, exemplified are fluoromethyl group, methyl-chloride group, bromomethyl group, and lodomethyl group.

[0105] Proferable substituents as a substituent capable of condensation polymerization depend upon a kind of polymerization reaction. For example, when using a zero-valent nickeckel complex, such as Yamamoto coupling reaction, exemplified are a halogen atom, alkly sulfonate group, any sulfonate group, or aylalky sulfonate group, Vene using a nickel catalyst or a palladium catalyst, such as Suzuki coupling reaction, exemplified are an alkyl sulfonate group,

halogen atom, boric ester group, boric axid group, etc.

[1016] When the polymer compound of the present invention has a repeating units other than a thiophene condensed
ring unit, a condensation polymerization can be carried out under the existence of a compound having two substituents
capable of condensation polymerization as the representing unit other than the thiophene condensed ring unit.

25 [0107] As the compound having two substituents capable of condensation polymerization used as a repeating unit other than formula (1-1) and (1-2), exemplified are the compounds of above formula (16)- (19).

$$Y_e\{Ar_0 - X_1\}_{11} - Ar_0 - Y_e$$
 (17)

$$Y_0 - X_0 - Y_{+0}$$
 (19)

wherein, Ar, A_{2} , A_{2} , A_{1} , B_{1} , A_{2} , B_{1} , A_{2} , and A_{3} are the same as the above. Y_{3} , Y_{4} , Y_{5} , Y_{8} , Y_{7} , Y_{8} , Y_{9} , A_{1} , and Y_{10} each independently represent a substituent capable of condensation polymerization. [0108] As the compound having two substituents capable of condensation polymerization used as a repeating unit other than formula (1-1) and (1-2), the compound represented by the below formula (2-1) and (1-2-1) performable.

$$Y_{27}$$
— Ar_{8} — N — Ar_{7} — N — X
 Ar_{8} — Y_{28}
 Ar_{10}
 N — Ar_{11}
 Ar_{12}

(24-1)

wherein, Ar₆, Ar₇, Ar₈, Ar₉, Ar₁₀, Ar₁₁, Ar₁₂, x, and y are the same as the above. Y₂₇ and Y₂₈ each independently represent a substituent capable of condensation polymerization.

[0109] A compound represented by formula (24-2) is further preferable.

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$$Y_{23} \underbrace{\begin{pmatrix} R_{50} \\ H_{51} \end{pmatrix}_{ii}}_{Z} Y_{30}$$

24-2)

wherein, R₅₀, R₅₁, R₅₂, hh, ii, jj, and z are the same as the above. Y₂₉ and Y₃₀ each independently represent a substituent capable of condensation polymerization.

[0110] Specifically, a compound used as a monomer, having a plurality of reactive substituents is dissolved, if necseary, in an organic solvent, and can be reacted at the melting empenture or more and the bolling polit or less of the organic selvent using an alkali or suitable catalyst, for example. For example, known methods can be used, described in "Organic Reactions", vol. 14, pp. 270 to 490, John Wiley & Sons, Inc., 1985, "Organic Reactions", vol. 27, pp. 345 to 390, John Wiley & Sons, Inc., 1982, "Organic Synthesis", Collective Volume VI, pp. 407 to 411, John Wiley & Sons, Inc., 1988, Chemical Review, vol. 95, p. 2457 (1995), Journal of Organicmetalic Chemistry, vol. 576, p. 147 (1998), Journal of Praticial Chemistry, vol. 336, p. 247 (1994), Makromolecular Chemistry Macromolecular Symposium, vol. 12, p. 229 (1987), and the like Vision of the Vision of Vision

[0111] In a method of producing the polymer compound of the present invention, as the method of carrying out condensation polymerization, a known condensation reaction can be used according to the substituent of the compound represented by the above formula (14) - (24-2) capable of condensation polymerization.

[0112] When the polymer compound of the present invention generates a double bond in the condensation polymer actization, a method recited in JPS 2-602565A is exemplified. Namely, exemplified are: a polymerization by Willig reaction of a compound having formyl group and a compound having phosphonitum methyl group, a polymerization by Heck reaction of a compound having singly group; a polymerization by Heck reaction of a compound having vinyl group and a compound having halogen atom; a polycondensation by delyational possible and a compound having to or more mon-halogenated methyl group; a polycondensation by the sutfontum-said decomposition method of a compound having when or more sutfontum-methyl groups; a polymerization by Knoevenagel reaction of a compound having a formyl group and a compound having down or more formyl groups, a formyl group and a compound having to or more formyl groups, at compound having to or more formyl groups, at compound having a formyl group and a compound having down or more formyl groups, at compound having a formyl group and a compound having town or more formyl groups, at compound having a formyl group and a compound having a formyl group and

[0113] When a polymer compound of the present invention generates a triple bond in the main chain in condensation polymerization, for example, Heck reaction can be used.

[0114] Moreover, when neither a double bond nor a triple bond is generated, exemplified are: a polymerizing method of a corresponding monomer by Suzuki coupling reaction; a polymerizing method by Grignard reaction; a polymerizing method by nickel (0) complex; a polymerizing method by nickel (0) complex; a polymerizing method by an oxidizing agent such as FeCl₃; an electrochemical oxidation polymerization method; or a method by decomposition of an intermediate polymer having a suitable leaving group.

[0115] Of these, the polymerization method by a Wittig reaction, the polymerization method by a freet ceaction, the polymerization method by a Knoevenagel reaction, the polymerization method by a Suzuki coupting reaction, the polymerization method by a Grignard reaction and the polymerization method using a NI(0) catalyst are preferable since structure control is easy in these methods.

[0116] In the manufacture methods of the present invention, it is preferable is that $Y_1, Y_2, Y_3, Y_4, Y_5, Y_5, Y_7, Y_8, Y_9$, and Y_{10} are each independently a halogen atom, alkyl sulfonate group, anyl sulfonate group, or anylalkyl sulfonate group, and condensation polymerization is carried out under the existence of zerovalent nickel complex.

[0117] As the raw material compounds, a dihalogenated compound, bis (alkylsulfonate)compound, bis(arylsulfonate) compound, bis (arylsulfonate)compound, bis (arylsulfonate)compound, bis (arylsulfonate)compound, bis (arylsulfonate)compound, bis (arylsulfonate)compound, bis (arylsulfonate)compound, alkylsulfonate compound, alkylsulfonate compound,

pound are exemplified.

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[0118] Moreover, in the manufacture method of the present invention, it is preferable that Y₁, Y₂, Y₃, Y₄, Y₅, Y₆, Y₇, Y₈, Y₈, Y₈, and Y₁₀ are each independently a halogen atom, alkyl sufforate group, anysulfonate group, anylarlysistionate group, borice adid group or boric eater group, and and the ratio of the total moles (0) of a halogen atom, alkylsulfonate group, anyl sufforate group, and anylarlysisulfonate group, to the total moles (K) of the boric acid group and boric ester group is substantially 1 (KV) is usually in the range of 0.7–1.2), and condensation polymerization is carried out using a nicleot catalyst or a palladium catalyst.

- [0119] As the concrete combination of the raw material compounds, exemplified is a combination of dihalogenated compound, bis[ally/sulfonate]compound, a bis (arysulfonate)compound or bis[aryslally/sulfonate]compound, with diboric acid compound or diboric ester compound.
- [0120] Moreover, exemplified are halogen-boric acid compound, halogen-boric ester compound, alkylsulfonate-boric acid compound, alkylsulfonate-boric ester compound, anylsulfonate-boric acid compound.
- 5 (1411) It is preferable that the organic solvent used is subjected to a deoxygenation treatment sufficiently and the reaction is progressed under an inent atmosphere, generally for suppressing a side reaction, though the treatment differs depending on compounds and reactions used. Further, it is preferable to conduct a dehydration treatment likewise (however, this is not applicable in the case of a reaction in a two-phase system with water, such as a Suzukl counting reaction).
- 20 [0122] For the reaction, an alkali or suitable catalyst is added appropriately. These may be selected according to the reaction used. It is preferable that the alkali or catalyst is soluble sufficiently in a solvent used for the reaction. As the method of mixing an alkali or catalyst, there is exemplified a method of adding a solution of an alkalist or catalyst slowly while stirring under an inner atmosphere of argon and nitrogen and the like or a method of slowly adding the reaction solution to a solution of an alkalia or catalyst. Inversely.
- 25 [0123] When these polymer compounds are used as a light-emitting material of a polymer LED, the purity thereof exerts an influence on light emitting ropperty, therefore, it is preferable that a monomer is purified by a method such as distillation, sublimation purification, recrystalization and the like before being polymerized and further, it is preferable to conduct a purification treatment such as re-precipitation purification, chromatographic separation and the like after the a varieties.
- 39 [0124] In case of X=S, the compound represented by the above formula (14) can be produced by reducing the compounds of below formula (22) or (23), with using a reducing agent.

wherein, V represents a repeating unit having the structure of X-SO₂ corresponding to the above U. W represents a repeating unit having the structure of X-SO corresponding to the above U. Y, and Y₂ are the same as above-mentioned. [0128] For example, the dibenzothiophene derivative represented by the above formula (15-1) can be produced by reducing the compound represented by the below formula (15-12) with sings a reducing agent.



Symptonic, Y₁₁, Y₁₂, B₃₆, B₃₆, aa, and bb are the same as above-mentioned.
[0126] The reducing agent is not specially limited, and examples thereof include, titanium compounds, such as titanium trichloride; silane compounds, such as tribolroosilane; aluminium hydride compounds, such as aluminium hydride.

- d- i-buyl aluminium hydride, lithium aluminium hydride, Irriventhoxy aluminium lithium hydride, Juri-butoxy aluminium olithium hydride, aluminium sodium hydride, biethy-may dainimium sodium hydride, biethy-may burinium sodium hydride, diethy-latinium sodium bydride sib-antely-te-ty-tyborane, disposite hey-tyborane, disposite hey-tyborane, disposite hydride, diethy-tyborane, disposite hey-tyborane, disposite hydride syanide, sodium borohydride syanide, tybutyboran sodium hydride, sodium borohydride, timeth-oxyboran sodium hydride, and sodium borohydride sodium borohydride sodium borohydride, sodiu
- [0128] The reduction reaction can be carried out in a solvent under an inert atmosphere, such as nitrogen or argon.

 The temperature of the reaction is preferably from -80°C to a boiling point of the solvent.
- [0128] As the solvents, exemplified are: saturated hydrocarbons, such as pentane, hoxane, heptane, octane, and cyclohexane; unsaturated hydrocarbons, such as benzene, toluene, ethylbenzene, and xylene; halogenated saturated hydrocarbons, such as carbon tetrachioride, chiorotomi, dichioromethane, chiorobetane, bromoperatane, bromoperatane, chiorobetane, and tromopolotokarue, halogenated unsaturated hydrocarbons, such as schioroberane, edichioroberane, and trichioroberane, alcohoroberane, alcohoroberane, and trichioroberane, and such such as formic acid, such as sformic acid, ethers, such as dimethylether, diethylisther, methyl-t-bully ether, tetrahydroturan, and doxane; and continues, such as the such such as the such
- [0130] As the post-treatment, a usual method is available, for example, quenching with water, and extracting by an 25 organic solvent, and then distilling the solvent. Isolation and purification of the product can be performed by methods such as factionation by chromatography, or convisitilization.
 - [0131] Next, the use of the polymer compound of the present invention is explained.
- [0132] The polymer compound of the present invention has fluorescence or phosphorescence in the solid state, and it can be used as a light emitting polymer (high molocular weight light-emitting material). Moreover, said polymer compound has excellent electronic transportation ability, and can be used suitably as a polymer-LED material, or a charge transporting material. The polymer LED using this light emitting polymer is a high performance polymer LED which can be driven at a low-voltage and efficiently. Therefore, the polymer LED can be preferably used for beak light of a liquid drystal display, a light source of curved or flat surface for lighting, a segment type display element, and apparatus such as a flat-panel display of oth matrix.
- [518] Moreover, the polymer compound of the present invention can be used also as a coloring matter for lasers, a material for organic solar-cell, an organic transistor for organic semiconductor, and a conductive thin film such as conductive thin-film material, and organic-semiconductor thin film.
 - [0134] Furthermore, it can be used also as a luminescent thin-film material which emits fluorescence or phosphorescence.
- 40 [0135] Next, the polymer LED of the present invention is explained.
 - [0138] The polymer LED of the present invention comprises an organic layer between the electrodes consisting of an anode and a cathode, and the organic layer contains the polymer compound of the present invention. The organic layer may be any of a light entiting layer, a hole transporting layer, and an electron transporting layer, and it is preferable that the organic layer is a light metting layer.
- 49 [0137] The light emitting layer means a layer having a function of light-emission. The hole transportating layer means a layer having a function of transporting layer means a layer having a function of transporting electrons. The electron transporting layer and the hole transporting layer are collectively referred to as a charge transport layer. Two or more layers of the light emitting layer, hole transporting layer, and electron transporting layer and the business of the light emitting layer, hole transporting layer, and electron transporting layer and electron transporting layer.
- [0138] When an organic layer is the light emitting layer, said light emitting layer which is an organic layer may contain further a hole transporting material, an electron transporting material, or a light-emitting material.
- [0138] The composition comprising the polymer compound of the present invention and at least one compounds as a light-emitting material and legit-emitting material can be used as a light-emitting material and legit-emitting material can be used as a light-emitting material or a charge transporting material. When mixing the polymer compound of the present invention and a hole transporting material, the mixed ratio of the hole transporting material is 1 wt%-80 wt% based on the total amount of the mixture, and it is preferably 5 wt%-80-bwt%. When mixing the polymer compound of the present invention and an electron transporting material, the mixed ratio of the electron transporting material is 1 wt%-80 wt%. But with the column of the mixture, and it is preferably 5 wt%-60-bwt%. Furthermore, when mixing the polymer

compound of the present invention and a light-emitting material, the mixed ratio of the light-emitting material is 1 wt% - 90 wt% based on the total amount of the mixture, and it is preferably 5 wt% - 60wt%.

(0140) When mixing the polymer compound of the present invention with a light-emitting material, a hole transporting material, and/or an electron transporting material, the mixed ratio of the light-emitting material is 1 wt% - 50 wt% based on the total amount of the mixture, and it is preferably 5 wt% - 40 wt%. The total amount of the hole transporting material and the electron transporting material and the present invention is 99wt%. 20mt%.

[0141] As the hole transporting material, electron transporting material, and fight-emitting material to bomixed, known town includes weight compounds, known phosphorescent compounds and known polymer compounds, gas an be used, and it is preferable to use a polymer compound. See the hole transporting material or copyimer compound, seem to compound the preferable to use a polymer compound, seem the preferable to use a polymer compound, seem the transporting material or copyimer compound, seem the rest of and copyimer shereof, a polyarylene winylene and derivatives thereof and copyimers thereof, a polyarylene winylene and derivatives thereof and copyimers thereof, a polyarylene winylene and derivatives thereof and copyimers thereof, and copyimers thereof and copyimers thereof, and copyimers ther

[0142] As the light-emitting material of low molecular weight compound, exemplified are: naphthalene, and derivatives thereof; anthracene and derivatives thereof, and derivatives thereof; coloring matters, such as polymethines, xanthenes, coumarins, and cyanines; 8-hydroxyquinoline or the metal complex of its derivative; aromatic amines; telrapheny occlopentadienes or its derivative; tetrapheny/butadiene or its derivatives, etc.

[0143] Specifically, known materials disclosed in JP 57-51781A or JP 59-194393A, can be used.

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[0144] As the phosphorescent compounds, exemplified are: those having iridium as a central metal, such as Ir(ppy) 3 and Btp_Ir(lacac); those having platinum as a central metal, such as PtOEP; those having europium as a central metal such as EurTHA)sohen, etc.

$$C_2H_5$$
 C_2H_5 C_2H_6 C_2H_6 C_2H_6 C_2H_6 C_2H_6 C_2H_6 C_2H_6 C_2H_6 C_2H_6 C_2H_6

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$$\begin{bmatrix} S \\ O \\ O \\ CF_3 \end{bmatrix}, Eu(TTA)_3 phet$$

[0145] Specifically, examples of the phosphorescent compounds are described in: Nature, (1998), 386, 151; App. Phys. Latt. (1999), 761(4), Fire SPIE-Inf. Soc. Ozt. Eng. (2001), 4106 (Organic Light-Hintilly) Atlariate and Devices IV), 119; J. Am. Chem. Soc., (2001), 123, 4304; Appl. Phys. Lett., (1997), 71(18), 2596; Syn. Met., (1998), 94(1), 103; Syn. Met., (1999), 992(1), 3817, 4407, Matter, (1999), 11(10), 882; Jun. J.Appl. Phys. 34, 1883 (1995), etc.

[0146] As a polymer compound to be mixed with the phosphorescent compounds, those represented by formula (1-1) having only the structure of X=S are preferable.

- [0147] Regarding the thickness of the light emitting layer, the optimum value differs depending on material used, and may properly be selected so that the driving voltage and the light emitting efficiency become an optimum value, and the thickness is, for example, from 1 mn to 1 um, preferably from 2 mn to 500 mn, further preferably from 5 mn to 200 mn.
 [0148] As a forming method of the light emitting layer, for example, exemplified is a method by film forming from a solution
- Id (14) As the film-forming method from a solution, application methods, such as a spin cost method, casting method, did not gravure coating method, par-coating method, roll coating method, which are coating method, solven printing, flexography method, offset printing, and lnk jet printing method, can be used. Printing method, such as screen printing, flexography method, offset printing, and ink jet printing method, are preferable, since pattern forming and multicolored printing are offsetable, since pattern forming and multicolored printing are offsetable.
- (0150) As the ink composition used for the printing method etc., at least 1 kind of the polymer compounds of the present invention should be contained, additives, such as a hole transporting material, electron transporting material, light-emitting material, solvent, or stabilizer, may be contained in addition to the polymer compound of the present invention.
- [0151] The amount of the polymer compound of the present invention in said ink composition is 20wt% 100wt% based on the total weight of the composition except the solvent, and preferably 40wt% 100wt%.
 - [0152] When a solvent is contained, the amount of the solvent in the ink composition is 1wt%-99.9wt% based on the total weight of the composition, preferably 60wt%-99.5wt%. and more preferably 80wt%-99.0wt%.
- [0153] The suitable viscosity of the ink composition is depend on the printing methods. When the ink composition is processed via a discharging appearatus, in order to prevent clogging and ejection-deflecting at discharging, the viscosity is preferably in a rance of 1 - 20mPa-s at 25°C.
- [0154] The solvent used as the ink composition is not especially limited, and preferable are those which can dissolve or disperse the materials constituting the ink composition other than the solvent uniformly.
- [0155] When the materials constituting the ink composition are soluble in a nonpolar solvent, as the solvent, exemplified are: chlorinated solvents, such as chiroromy, methylene chloride, and dichlorothane; other solvents, such as tatrihydrofurar; aromatic hydrocarbon solvents, such as total and yviene; ketone solvents, such as acottone, and methy othyr ketone control and solvents, such as a total solvents, such as acottone, and methy othyr ketone; and destroyents, such as are vital sociate, but will carelate, and their collections of acotter collections.
- [0156] Moreover, as the polymer LED of the present invention, there are exemplified: a device having an electron transporting layer disposed between a cathode and a light emitting layer; a device having a hole transporting layer.

disposed between an anode and a light emitting layer; and a device having an electron transporting layer disposed between a cathode and a light emitting layer, and a hole transporting layer disposed between an anode and a light emitting layer.

[0157] For example, the following structures a) to d) are specifically exemplified.

a) anode/light emitting laver/cathode

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- b) anode/hole transporting layer/light emitting layer/cathode
- c) anode/light emitting layer/electron transporting layer/ /cathode
- d) anode/hole transporting layer/light emitting layer/electron transporting layer/cathode
- (wherein, / indicates adjacent lamination of layers. Hereinafter, the same)

[0158] When the polymer LED of the present invention has a hole transporting layer, as the hole transporting materials used, there are exemplified polymicplactures or derivatives thereof, polyalizen or not enviratives thereof, polyalizen or not hoyalizen and enviratives that a remain that is a size of the main chain, purzoziline derivatives, any jamine derivatives, polyamilizen of enviratives thereof, polygorie or derivatives, triphenyfoliamine derivatives, polyamiline or derivatives thereof, poly(2,5-thienyfonevirnyfone) or derivatives thereof, poly(2,5-thienyfonevirnyfone) or derivatives thereof, poly(2,5-thienyfonevirnyfone) or derivatives thereof.

[0159] Specific examples of the hole transporting material include those described in JP 63-70257A, JP 63-175860A, JP 2-135359A, JP 2-135361A, JP 2-209988A, JP 3-37992A and JP 3-152184A.

[0160]. Among them, as the hole transporting materials used in the hole transporting layer, preferable are polymer hole transporting materials used as polymyined tables are polymer thereof, polysisism or derivatives having an aromatic amine compound group in the side chain or the main chain, polyanilitie or delocation derivatives having an aromatic amine compound group in the side chain or the main chain, polyanilitie or derivatives thereof, polythispheme or derivatives thereof, polythispheme yor derivatives thereof, polythispheme yor derivatives thereof, polythispheme are polythispheme or derivatives thereof, polythispheme with preferable are polythispheme or derivatives thereof and obstitutionare derivatives having an aromatic amine compound group in

the side chain or the main chain.

(01611 Moreover, as the hole transporting material of low molecular weight compound, exemplified are pyrazoline

derivatives, anylamine derivatives, stilbene derivatives, and triphenyl diamine derivatives. In case of the low molecular weight hole transporting material, it is preferably dispersed in a polymer binder for use.

90 [0162] As the polymer binder to be mixed, pelerable are those which do not high thich darge transportation extremely and do not have strong absorbance in a visible light. As the polymer binder, exempiffled are; poly (N-vinylcarbazole);

polyaniline or derivatives thereof; poly thiophene or derivatives thereof; poly(p-phenylynenvirylene) or derivatives thereof; of; poly(2,5-thienylenevirylene) or derivatives thereof; polyeabonate; polyacrylate, poly methylacrylate, polymethylacrylate, polyetyrene, polyvinylchloride, polyslozane, etc. (0163) Polyvinyl carbazole and derivatives thereof are obtained from, for example, a vinyl monomer, by cationic

polymerization or radical polymerization.

[0164] As the polysilane or derivalives thereof, there are exemplified compounds described in Chem. Rev., 89, 1359 (1989) and GB 2300196 published specification, and the like. For synthesis, methods described in them can be used, and a Kipoing method can be suitably used particularly.

40 [0165] As the polysiloxane or derivatives thereof, those having the structure of the above-described hole transporting material having lower molecular weight in the side chain or main chain, since the siloxane skeleton structure has poor hole transporting property. Particularly, there are exemplified those having an aromatic amine having hole transporting property in the side chain or main chain.

[0166] The method for forming a hole transporting layer is not restricted, and in the case of a hole transporting layer. A having lower molecular weight, a method in which the layer is formed from a mixed soution with a polymer binder is exemplified. In the case of a polymer hole transporting material, a method in which the layer is formed from a solution is exemplified.

[0167] The solvent used for the film forming from a solution is not particularly restricted providing it can dissolve a hole transporting material. As the solvent, there are exempfified chlorine solvents such as chloroform, methylene chloride, dichloroethane and the like, ether solvents such as tetrahydrofuran and the like, aromatic hydrocathon solvents such as toluene, xylene and the like, ketone solvents such as acetone, methyl ethyl ketone and the like, and ester solvents such as ethyl acetate, othyl acetate, ethyleoticsolve acetate and the like.

[0168] As the film forming method from a solution, there can be used coating methods such as a spin coating method, casting method, micro gravure coating method, pet coating method, relic casting method, micro gravure coating method, bet are called method, relic casting method, graving coating method, serven printing method, flexo printing method, graving method, offset printing method, flexible training method, misket printing method and the file. From a solution.

[0169] Regarding the thickness of the hole transporting layer, the optimum value differs depending on material used, and may properly be selected so that the driving voltage and the light emitting efficiency become an optimum value.

- The thickness is should be at least such that pinholes are not generated, but when it is too thick, the driving voltage of a device becomes high, and it is not preferable. Therefore, the fillin thickness of the hole transporting layer is, for example, from I mm to 1 um, preferably 2mm 500mm, and more preferably 5mm 500mm, and 5mm 200mm.
- [0170] When the polymer LED of the present invention has an electron transporting layer, known compounds are used as the electron transporting materials, and there are exemptified covalizable derivatives, anthraquinonel methane or derivatives thereof, benzoquinone or derivatives thereof, benzoquinone or derivatives thereof, intransparanthraquinodimethane or derivatives thereof, fluorence derivatives, deprendigatives, deprendigatives, deprendigatives of derivatives thereof, polyquinoline or derivatives thereof, polyquinoline and derivatives thereof, polyquinoline and derivatives thereof, polyquinoline and derivatives thereof, polyquinoline and derivatives.
- [0171] Specifically, there are exemplified those described in JP 63-70257A, JP 63-175860A, JP 2-135359A, JP 2-135351A, JP 2-20988A, JP 3-37992A and JP 3-152184A.
 - [0172] Among them, oxadiazole derivatives, benzoquinone or derivatives thereof, anthraquinone or derivatives thereof, or metal complexes of 8-hydroxyquinoline or derivatives thereof, polyquinoline and derivatives thereof are preferable, and 2-(4-biphenyl)-5-(4-bibyhphenyl)-1-3.4-oxadiazole, benzoquinone, anthraquinone, trisle-quinolinollatuminum and polyquinoline are further prefer-
- ny):1,:3,4-oxadiazose, benzoquinone, aninaquinone, inste-quinoiniopialuminum and polyquinoine are lutrier preterable.
 [0173] The method for forming the electron transporting layer is not particularly restricted, and in the case of an electron transporting material having lower molecular weight, a vapor deposition method from a powder, or a method
 - of film-forming from a solution or melled state is exemptified, and in the case of a polymer electron transporting material, a method of film-forming from a solution or melled state is exemptified, respectively. At the time of film forming from a solution or a mother state, the above polymer binder can be used together.

 [0174] The solvent used in the film-forming from a solution is not particularly restricted provided it can dissolve elec-
- tron transporting materials and/or polymer binders. As the solvent, there are exemplified chlorine solvents such as 25 chloroform, methylene chloride, dichlorochane and the like, either solvents such as tetrahydroturan and the like, aromatic hydrocarbon solvents such as toluene, xylene and the like, ketone solvents such as acetone, methyl ethyl ketone and the like, and ester solvents such as ethyl acetate, buyl acetate, ethylcollosolve acetate and the like.
- [0175] As the film-forming method from a solution or melted state, there can be used coating methods such as a spin coating method, casting method, micro gravure coating method, gravure coating method, ser coating method, and o coating method, wire bar coating method, dip coating method, spray coating method, screen printing method and the like.
- [0178] Regarding the thickness of the electron transporting layer, the optimum value differs depending on material used, and may properly be selected so that the driving voltage and the light emitting efficiency become an optimum value. The thickness is should be at least such that pinholes are not generated, but when it is too thick, the driving ovoltage of a device becomes high, and it is not preferable. Therefore, the film thickness of the hole transporting layer is, for example, from 1 min to 1, my preferably 5 mm. 500mm, and more perferably 5 mm. 200mm.
 - [0177] Among charge transporting layers provided adjacent to an electrode, those having a function to improve the charge linjection efficiency from an electrode and to lower the driving voltage of a device may be sometimes called charge injection layers (a hold-injection layer, electronic injection layer).
- 40 [0178] For enhancing adherence with an electrode and improving charge injection from an electrode, the above-described charge injecting layer or insulation layer having a thickness of 2 mm or less may also be provided adjacent to an electrode, and further, for enhancing adherence of the interface, preventing mixing and the like, a thin buffer layer may also be inserted into the interface of a charge transporting layer and light emitting layer.
- [0179] The order and number of layers laminated and the thickness of each layer can be appropriately applied while considering light emitting efficiency and life of the device.
 - [0180] In the present invention, as the polymer LED having a charge injecting layer (electron injecting layer, hole injecting layer) provided, there are listed a polymer LED having a charge injecting layer provided adjacent to a cathode and a polymer LED having a charge injecting layer dialecent to an anode.
- [0181] For example, the following structures e) to p) are specifically exemplified.
 - e) anode/charge injecting layer/light emitting layer/cathode

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- f) anode/light emitting layer/charge injecting layer/cathode
- g) anode/charge injecting layer/light emitting layer/charge injecting layer/cathode
- h) anode/charge injecting layer/hole transporting layer/light emitting layer/cathode
- i) anode/hole transporting layer/light emitting layer/charge injecting layer/cathode
- i) anode/charge injecting layer/hole transporting layer/light emitting layer/charge injecting layer/cathode
- k) anode/charge injecting layer/light emitting layer/electron transporting layer/cathode
- I) anode/light emitting layer/electron transporting layer/charge injecting layer/cathode

m) anode/charge injecting layer/light emitting layer/electron transporting layer/charge injecting layer/cathode n) anode/charge injecting layer/light emitting layer/elight emitting layer/electron transporting layer/cathode o) anode/hole transporting layer/light emitting layer/electron transporting layer/elayer injecting layer/elathode p) anode/charge injecting layer/hole transporting layer/light emitting layer/electron transporting layer/charge injecting layer/elathode

[0182] As the specific examples of the charge injecting layer, there are exemplified: layer containing an anotholding ophymor; layers which are disposed between an another and another anoth

[0183] When the above-described charge injecting layer is a layer containing an conducting polymer, the electric conductivity of the conducting polymer is preferably 10⁻⁵ S/cm or more and 10⁵ S/cm or less, and for decreasing the leak current between light entities pixels, more preferably 10⁻⁵ S/cm or more and 10² S/cm or less, further preferably 10⁻⁵ S/cm or more and 10³ S/cm or less.

9 (1941) Regarding the kind of an ion doped, an anion is used in a hole injecting layer and a cation is used in an electron injecting layer. As examples of the anion, a polystyness usefunate ion, allytherpacene sulfonate ion, camphor sulfonate ion and the like are exemplified, and as examples of the cation, a lithium ion, sodium ion, potassium ion, tetrabuly ammonium ion and the like are exemplified.

The thickness of the charge injecting layer is for example, from 1 nm to 100 nm, preferably from 2 nm to 50 nm.

20 [0185] Specifically, there are listed the following structures q) to ab) for example.

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q) anode/insulation layer having a thickness of 2 nm or less/light emitting layer/cathode r) anode/light emitting layer/insulation layer having a thickness of 2 nm or less/cathode

s) anode/insulation layer having a thickness of 2 nm or less/light emitting layer/insulation layer having a thickness of 2 nm or less/cathode

1) anodórisulation layer having a thickness of 2 mm or less/hole transporting layer/light emitting layer/cathode u) anode/hole transporting layer/flor emitting layer/mulation layer having a thickness of 2 mm or loss/cathode u) anodérisulation layer having a thickness of 2 mm or loss/hole transporting layer/flight emitting layer/flight and transporting layer/flight emitting layer/flight emit

30 w) anode/insulation layer having a thickness of 2 mm or less/light entiting layer/selectron transporting layer/scathode y anode/injet entiting layer/selectron transporting layer/insulation layer having a thickness of 2 mm or less/scathode y) anode/insulation layer having a thickness of 2 mm or less/scathode

anode/insulation layer having a thickness of 2 nm or less/hole transporting layer/light emitting layer/electron transporting layer/cathode

 aa) anode/hole transporting layer/light emitting layer/electron transporting layer/insulation layer having a thickness of 2 nm or less/cathode

ab) anode/insulation layer having a thickness of 2 nm or less/hole transporting layer/light emitting layer/electron transporting layer/insulation layer having a thickness of 2 nm or less/cathode

[0188] The substrate forming the polymer LED of the present invention may preferably be that does not change in forming an electrode and layers of organic materials, and there are exemplified glass, platetics, polymer film, silicon substrates and the like. In the case of a opaque substrate, it is preferable that the opposite electrode is transparent or semitransparent.

45 [0187] Úsually, either the anode or the cathode in the polymer LED of the present invention is transparent or semitransparent. It is preferable that the anode side is transparent or semitransparent.

[0188] As the material of this anode, electron conductive metal oxide films, semitransparent metal thin films and the like are used. Spocifically, there are used influm oxide, zinc oxide, into oxide, and films (NESA and the like) fabricated by using an electron conductive glass composed of Indium-tin-oxide (ITO), indium-zinc-oxide and the like, which are metal oxide complexes, and gold, platinum, silver, copper and the like are used, and among them, ITO, indium-zinc-oxide, in oxide are preferable. As the fabricating method, a vacuum vapor deposition method, sputtering method, on plating method, plating method and the like are used. As the anode, there may also be used organic transperent conducting films such as polyanilino or ordervalievs thereot, polythiophene or derivatives thereof and the like.

[0189] The thickness of the anode can be appropriately selected while considering transmission of a light and electric so conductivity, and for example, from 10 nm to 10 μ m, preferably from 20 nm to 1 μm, further preferably from 50 nm to 500 nm.

[0190] Further, for easy charge injection, there may be provided on the anode a layer comprising a phthalocyanine derivative conducting polymers, carbon and the like, or a layer having an average film thickness of 2 nm or less com-

prising a metal oxide, metal fluoride, organic insulating material and the like.

[0191] As the material of a cathode used in the polymer LED of the present invention, that having lower work function is preferable. For example, there are used metals such as lithium, sodium, potassium, nubridum, cesium, berjlüum, magnesium, calcium, stontium, barium, aluminum, scandium, vanedium, zinc, yftrum, indium, cerium, semarium, europium, terbium, terbium and the like, or alloys comprising two of more of them, or alloys comprising one or more of them with no en or more of gold, silver, platium, copper, managenes, trainium, coalair, lockel, fungstein antilin, graphite or graphite intercalation compounds and the like. Examples of alloys include a magnesium-aliver alloy, magnesium-indium alloy, magnesium-aluminum alloy, indium-either alloy, thithum-aluminum alloy, indium-either alloy, thithum-aluminum alloy, thit of the cathode may be formed tind a laminated structure of two or

10 more layers.
[0192] The thickness of the cathode can be appropriately selected while considering transmission of a light and electric conductivity, and for example, from 10 nm to 10 μm, preferably from 20 nm to 1 μm, further preferably from 20 nm to 1 μm, further preferably from 20 nm to 1 μm.

[0193] As the method for fabricating a cathode, then are used a vacuum vapor deposition method, sputtering method, iamination method in which a metal thin film is adhered under heat and pressure, and the like. Further, there may also be provided, between a cathode and an organic layer, a layer comprising an conducting polymer, or a layer having an average film thickness of 2 mm or less comprising a metal oxide, metal fluoride, organic insulation material and the like, and after radication of the cathode, a protective layer may also be provided which protects the polymer LED. For stable use of the polymer LED for a long period of time, it is preferable to provide a protective layer and/or protective over for protection of the device in order to prevent it from oxidised damage.

Ordan placeutori on the device in ordan to prevent in front outsale damage.

(1944) As the protective layer, there can be used a polymer compound, metal oxide, metal fluoride, metal borate and the like. As the protective cover, there can be used a glass plate, a plastic plate the surface of which has been subjected to lower-water-permetion treatment, and the like, and there is sublety used a method in which the cover is pasted with an device substrate by a thermosetting resin or light-curing resin for sealing. If space is maintained using a spacer, it is easy to prevent an device from being injured. If an inner gas such as nitrogen and argon is sealed in this space, it is possible to prevent oxidation of a cathode, and utture, by placing a desiccant such as barium oxide and the like in the above-described space, it is easy to suppress the demage of an device by moisture adhered in the production process. Among them, are none means or more are preferably adopted them.

[0195] The polymer LED of the present invention can be suitably used as a flat light source, segment display apparatus, dot-matrix display apparatus, and back light of a liquid crystal display.

(0198) For obtaining light emission in plane form using the polymer LEO of the present invention, an anode and a cathode in the plane form may properly be placed so that they are suminated each other. Further, for obtaining light emission in pattern form, there are a method in which a mask with a window in pattern form is placed on the above-described plane light emitting device, a method in which an organic layer in non-light emission part is formed to obtain a strength large thickness providing substantial non-light emission, and a method in which any one of an anode or a cathode, or both of them are formed in the pattern. By forming a pattern by any of these emitodes and by jacing some electrodes so that independent or/off is possible, there is obtained a display device of segment type which can display digits, laters, simple marks and the like. Further, for forming a dot matrix device, it may be advantageous that anodes and cathodes are made in the form of stripes and placed so that they cross at right angies. By a method in which a color filter or luminescence converting filter is used, area color displays and multi color displays are obtained. A dot matrix display can be driven by passive driving, or by active driving combined with TTF and the like. These display devices can be used as a display of a computer, television, portable terminal, portable telephone, car navigation, view finder of a video camera, and the like.

© [0197] Further, the above-described light emitting device in plane form is a thin self-light- emitting one, and can be suitably used as a flat light source for beck-light of a liquid crystal display, or as a flat light source for illumination. Further, if a flexible plate is used, it can also be used as a curved light source or a display.

EXAMPLES

[0198] The following examples further illustrate the present invention in detail but do not limit the scope theories (0199) As for he molecular weight, a number-everage molecular weight and a weight-everage molecular weight were obtained as a polystyrene reduced average molecular weight by gel permeation chromatography (GPC) using chioroform as a solvent.

Synthetic Example 1 (synthesis of Compound A)

[0200]

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HO ()

Compound A

[0201] 2.8-dibromodibenzothiophene 7g and THF 280ml were charged into a 1 liter four-necked flask under an inert atmosphere, and stirred to dissolve at a rown temperature, then cooled to 7% C. n-bullhitmian 28ml (1 Smok). havane solution) was added dropwise to it. After the dropwise addition, it was stirred for 2 hours, at the fixed temperature, and trimethoxyboronic acid if 3g was added dropwise. After the dropwise addition, the temperature was raised to a room temperature (swort), After stirring for 5 hours at room temperature, dissepperature of the rematerial was confilmed by TLC. 5% sulfuric acid 100ml was added to terminate the reaction, and stirred at a room temperature for 12 hours. After washing with water, the organic layer was extracted. After replacing the solvent with eithy acetals, 30% auguous hydrogen peroxide firm was added and stirred at 40°C for 5 hours. Then the organic layer was extracted, washed with 10% auguous outlion of irroll/gammonium sulfate, and dried. by removing the solvent, brown solid 4.43 was obtained.

By products, such as a dimer, were also produced as confirmed by LC-MS measurement, and the purity of Compound A was 77% (LC care precentage).

25 MS(APCI(-)):(M-H)* 215

Synthetic Example 2 (synthesis of Compound B)

[0202]

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Compound B

40 [2033] Under an inert atmosphere, into a 200ml three neck flask, Compound A.4.3g, n-octylpromide 25.1 g and potassium carbonate 12.5g were charged, and methylsocutyl ketone 50ml was added as a evolvent, then refluxed with heating at 128°C for 8 hours. After the reaction, the solvent was removed, and the reaction product was separated with chioroform and water, the organic layer was extracted and further washed with water twice. After driply by arityrous sodium suifate, it was purified through silica got column (eluent: toluene/cydohexane=1/10) to give Compound 45 8 8.49 fil. Care % = 97%, viol-94% was obtained.

<sup>1</sup>H-NMR(300MHz/CDCl<sub>3</sub>): δ 0.91 (t, 6H), 1.31-1.90 (m, 24H), 4.08 (t, 4H), 7.07 (dd, 2H), 7.55 (d, 2H), 7. 68 (d, 2H)

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Synthetic Example 3 (synthesis of Compound C)

[0204]

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Compound C

[0205] Compound B 6.67g and acetic acid 40ml were chargod into a 100ml three neck flask, and the bath-temperature was raised to 140 °C with using an oil bath. Then, 30% hydrogen-peroxide 13ml was added through a condenser, and attreed vigorously, then the reaction was terminated by being poured into 180ml of cold water. After attracting with chioroform and being dried, the solvent was removed and Compound C 6.99g (LC area % = 90%, yleid=97%) was obtained.

1H-NMR(300MHz/CDCl<sub>3</sub>):

δ 0.90 (t, 6H), 1.26-1.87 (m, 24H), 4.06 (t, 4H), 7.19 (dd, 2H), 7.69 (d, 2H), 7.84 (d, 2H) MS(APCI(+1):(M+H)+ 473

Synthetic Example 4 (synthesis of Compound D)

25 [0206]

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Compound D

[0207] Compound C 3.86g and mixed solution 15ml of acetic acid/ chloroform =1:1 were charged into a four-necked flask 20ml under an inert atmosphere, stried at 7 or C and dissolved. Then, bromine 6.02g dissolved in the above solvent 3ml was added and stirred for 3 hours. A sodium thiosulfate aqueous solution was added to remove unreacted bromine, and the reaction product was separated with chloroform and water, the organic keyer was extracted and dired. After removing the solvent, it was purified through silica get column (eluent: chloroform/hexane=1/4) to give Compound D 4.48g (LC area %= 98%, yield=84%) was obtained.

8 0.95 (t, 6H), 1.30~1.99 (m, 24H), 4.19 (t, 4H), 7.04 (s, 2H), 7.89 (s, 2H) MS(FD+)M+630

Example 1 (synthesis of Compound E)

[0208]

Compound E

[0209] Compound D 3.9g and diethyl either 50ml were charged into a 200ml three neck flask under an inent atmosphere, and the temperature was raised to 40°C and stirred. Lithium aluminium hydride 1.17g was added a title at a time and reacted for 5 hours. Excess of lithium aluminium hydride was decomposed by adding a small amount of water at a time, and washed with 50% hydrochloric acid 5.7ml. The reaction mixture was separated with chiloroform and water, the organic layer was extracted and dried. It was purified through silics get column (eluent: chloroform/hexane-15) to give Compound E 1.8g (LC area % =99%, yield=49%) was obtained.

δ 0.90 (t, 6H), 1.26~1.97 (m, 24H), 4.15 (t, 4H), 7.45 (s. 2H), 7.94 (s, 2H)

10 [0210] According to MS (APCI (+)) method, peaks were observed at 615 and 598.

Evample 2

<Synthesis of Polymer Compound 1>

[0211] After dissolving Compound E 362mg and 2,2'-bipyridyl 274mg in tetrahydrofuran (dehydrated) 20ml, the Inside of the system was replaced with nitrogen by bubbling with nitrogen gas. To this solution, bist(1,5-cyclooctadiene)nickel (0) (NI)(COD)<sub>2</sub> 500mg was added, and the temperature was raised to 50°C and reacted for 5 hours. After the reaction, this reaction liquid was cooled to a room temperature (about 25°C), and added dropwise to a mixed solution of 25% aqueous ammonia 10ml/ methanol 120ml/ 50ml in over-kanaged water and sturred for 1 hour. The deposited precipitate was fiftrated and dried for 2 hours under reduced pressure, and dissolved in toluene 30ml. Then, 30ml of 1 N hydrochloric acid was added and stilmed for 1 hour, the aqueous layer was removed, 30ml of 4% aqueous ammonia was added to the organic layer, and the equeous layer was removed after stirring for 1 hour. The organic layer was added dropwise to methanol 200ml, and stirred for 1 hour, the adject was present to the control of the contro

The polystyrene reduced average molecular weight of Polymer Compound 1 was Mn=2.1x104, and Mw=8.1x104.

Example 3

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<Synthesis of Polymer Compound 2>

35 (0212) A copolymer was produced according to the procedure of Example 2, with using Compound E 253mg, and N,N'-bis(4-bromophenyl)+N,N'-bis(4-broulybenryl)-1,4-phenylenedlemine 125mg as the starting materials.
(0213) Yield of the copolymer was 127mg, In the copolymer, the ratio of the repeating unit of disenzothicphene to the arrine derivative was 70:30. This copolymer is referred to as Polymer Compound 2. The polystyrene reduced average melecular weeldn of Polymer Compound 2 was Mne. 23.014, and Mne. 5s.xtf0-6.

Example 4

<Synthesis of Polymer Compound 3>

45 [0214] A copolymer was produced according to the procedure of Example 2, with using Compound E 296mg, and N,N-bis(4-bromopheny)<sup>1</sup>-N'N-diphenyl-1.4-phenylenediamine 144mg as the starting materials. [0215] 'Yield of the copolymer was 118mg, in the copolymer, the ratio of the repeating units of dibenzothiophene to the amine derivative was 70:30. This polystyrene reduced average molecular weight of Dyomer Compound 3 was Mn=2.6x16\*, and Mw=8.9x16\*.

Example 5

<Synthesis of Polymer Compound 4>

[0216] Compound E 6450mg, N.N-bis(4-bromophenyl)-N-(4-(1-methylbutyl) phenyl) amine 2070mg and 2,2'-bipyridyl 5500mg were charged into a flask, after the inside of the system was replaced with nitrogen, they were dissoved in 400mil of dhydrated tetrahydrofuran degassed with argon gas. To this solution, under nitrogen atmosphere, bis (1,5-cycloodadiene)nickel(0)(NI(COD)<sub>2</sub>) 1000mg was added, and the temperature was raised to 60°C and reacted

for 3 hours

After the reaction, this reaction mixture was cooled to a room temperature, and added dropwise to a mixed solution of 25% agueous ammonia 100ml methanol 500ml / 100ml lon-exchanged water and sitter for 1 hour. The deposited precipitate was filtrated and dried for 2 hours under reduced pressure, and dissolved in toluene 40ml. Then, 40ml of 1 N hydrochloric acid was added and stirred for 1 hour, the agueous layer was morrowed, 40ml of 3% aqueous sammonia was added to the organic layer, and the aqueous layer was removed after sittings for 1 hour. After washing with 150ml lon-exchanged water, the washed organic layer was poured into methanol 100ml, and stirred for 1 hour. Then deposited precipitate was littered and dried under reduced pressure for 2 hours, and dissolved in toluene 40ml. Then, purification through alumina column (25g of alumina) was carried but, and recovered toluene solution was added to methanol 200ml, stirred for 1 hour, and deposited precipitate was filtrated and dried under reduced pressure for two methanol 200ml, stirred for 1 hour, and deposited precipitate was filtrated and dried under reduced pressure for two methanol 200ml, stirred for 1 hour, and deposited precipitate was filtrated and dried under reduced pressure for two

[0217] Yield of the copolymer was 4000mg. In the copolymer, the ratio of the repeating units of dibenzothiophene derivative to the amine derivative was 70:30. This copolymer is referred to as Polymer Compound 4. The polystyrene reduced average molecular weight of Polymer Compound 4 was Mn=4.8x10\*, and Mw=5.xx10\*.

Example 6

<Synthesis of Polymer Compound 5>

- 20 [0218] Compound E 400mg, N.N'-biel,3'-methyl-4-bromophenyldiphenylbenzidine 440mg and 2,2'-bipyridyl 510mg were charged into a reaction flask, and the inside of the systems explaced with intogen. To this, 40mf of dehydrated tetrahydrofuran degassed by bubbling with argon gas was added. To this mature, biel (1,5-cyclocatedine)nickel(0) 1000mg was added, and a reaction was carried out at 80°C for 3 hours. The reaction was carried out under introgen atmosphere. After the reaction, this reaction mixture was cooled, and added dropwise to a mixed solution of 25% aqueous ammonia 100ml/ methanol 120ml/ 50ml ion-exchanged water and stirred for about 1 hour. The deposited precipitate was recovered by filtration. After being washed with eithanol, the precipitate was died of 27 hours under reduced pressure. Then the precipitate was died solved in toluene 30ml, then, 30ml of 1 N hydrochloric acid was added and stirred for 1 hour. The advocus layer was removed, 30ml of 14% aqueous ammonia was added to the organic layer, and dissolved in toluene 30ml. Then, puffication through alumina column (20g of alumina) was acarried out, and recovered toluene solution was added to the methanol 150ml, and stirred for 1 hour. The dissolved in toluene 30ml. Then, puffication through alumina column (20g of alumina) was acarried out, and recovered toluene solution was added to methanol 100ml, stirred of 1 hour, and deposited prepoblitate was filtrated and drift of the probelitate was filtrated and drift of the probelitate was filtrated and drift.
- [0219] Yield of the copolymer was 79mg. In the copolymer, the ratio of the repeating units of dibenzothiophene of derivative to the amine deviative was 50:50. This copolymer is referred to as Polymer Compound 5. The polystyrene reduced average molecular weight of Polymer Compound 5 was Mn=2.0x10<sup>5</sup>, and Mw=3.4x10<sup>4</sup>.

Example 7

40 <Synthesis of Polymer Compound 6>

under reduced pressure for two hours.

[0220] A copolymer was produced according to the procedure of Example 6, with using Compound E 400mg, and 2,3-diethyl-5, 8-dibromoquinoxaline 97mg.

[0221] Yield of the copolymer was 50mg. In the copolymer, the ratio of the repeating unit of dibenzothlophene dedrauler to quinoxaline unit was 70:30. This copolymer is referred to as Polymer Compound 6. The polystyrene reduced average molecular weight of Polymer Compound 6 was Mm-2,510f; and Mw-9,0x10f.

Example 8

<A synthesis of a polymer compound 7>

[0222] A copolymer was produced according to the procedure of Example 6, with using Compound E 350mg, NN\*bis(4-bromopheryl)+N,N\*-bis (4-n-butylphenyl) 1,4-phenylenediamine (amine derivative 1) 57 mg, and N,N-bis (4-bromophenyl)+N-(4-(1-methyl butyl) phenyl) amine (amine derivative 2) 78mg.

[0223] Yield of the copolymer was 76mg. In the copolymer, the ratio of the repeating unit of diborazothiophene derivative: amine derivative 1: amine derivative 2 was 70.10.20. This copolymer is referred to as Polymer Compound 7. The polystyrene reduced average molecular weight of Polymer Compound 7 was Mn=7.4410<sup>4</sup>, and Mw=2.4410<sup>5</sup>.

Example 9

<Synthesis of Polymer Compound 8>

5 (0224) A copolymer was produced according to the procedure of Example 5, with using Compound E 310mg, N, N-16 (4-bromopheny)-N, N-16-bit (4-bromopheny)

Example 10 (Synthesis of Compound F)

[0226]

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Br—Br

Compound F

25 Compound F was synthesized according to the procedure of the synthesis of Compound E from Compound A. Yield 46%, and purity 96%.

1H-NMR(300MHz/CDCl<sub>2</sub>):

δ 0.89 (dd, 12H), 1.01 (d, 6H), 1.17-2.01 (m, 20H), 4.18 (t, 4H), 7.42 (s, 2H) and 7.92 (s, 2H) According to a MS (APCI (+)) method, peaks were observed at 671 and 654.

Example 11

<Synthesis of Polymer Compound 9>

- 35 [0227] A polymer was synthesized according to the procedure of Example 5, using Compound F 1000mg. The yield of resultant polymer was 100mg. This polymer is referred to as Polymer Compound 9.
  [0228] The polystyren ecduced average molecular weight of Polymer Compound 9 was Mn=5.0x10<sup>4</sup>, and
  - [0228] The polystyrene reduced average molecular weight of Polymer Compound 9 was Mn=5.0x10°, and Mw=1.1×10<sup>5</sup>.
- 40 Example 12

<Synthesis of Polymer Compound 10>

- [0229] According to the procedure of Example 6, it was synthesized using Compound F 320mg and Compound E 35 310mg. In the copolymer, the ratio of the repeating units corresponding to Compound F and Compound E is 50:50. This polymer is referred to as Polymer Compound 10.
  - [0230] The polystyrene reduced average molecular weight of Polymer Compound 10 was Mn=8.8x104, and Mw=3.0x105.
- 50 Example 13

<Synthesis of Polymer Compound 11>

- [0231] According to the procedure of Example 6, it was synthesized using Compound F 320mg and N,N'-bis (4-bromophenyl)-N,N'-bis(4-n-butylphenyl)-1,4-phenylenediamine 140mg.
  - [0232] The yield of the resultant copolymer was 150mg. In the copolymer, the ratio of the repeating units of dibenzothiophene derivative and arrine derivative is 70:30.
  - [0233] This copolymer is referred to as Polymer Compound 11. The polystyrene reduced average molecular weight

of Polymer Compound 11 was Mn=3.9x103, and Mw=4.4x104.

Example 14

5 <Synthesis of Polymer Compound 12>

[0234] According to the procedure of Example 6, it was synthesized using compound F 270mg, compound E 250mg, and N, N-bis (4-bromophenyl)-N,N-bis (4-n-buylphenyl)-1,4-phenylenediamine 240mg, in the copoplamer, the ratio of the repeating units corresponding to Compound F. Compound E, and the amine derivative is \$5.35.30.

10 [0235] This copolymer is referred to as Polymer Compound 12. The polystyrene reduced average molecular weight of Polymer Compound 12 was Mn=3.8x10<sup>4</sup>, and Mw=5.9x10<sup>4</sup>.

Example 15 (Synthesis of Compound G)

15 [0236]

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Br Br

Compound G

[0237] It was synthesized according to the procedure of the synthesis of Compound E from compound A. 53% of yield, and 100% of purity.

14-NMR/300MHz / CDCiA:

δ 1.00 (d, 12H), 1.75 (dd, 2H), 1.87 (m, 4H) 4.11 (t, 4H), 7.07 (dd, 2H), 7.55 (d, 2H) and 7.68 (s, 2H)

Example 16

35 <Synthesis of Polymer Compound 13>

[0238] According to the procedure of Example 6, it was synthesized using Compound G 260mg and Compound E 300mg. The yield of the resultant polymer was 150mg. In the copolymer, the ratio of the repeating units corresponding to Compound 6 and Compound E is 6056.

40 [0239] This polymer is referred to as Polymer Compound 13. The polystyrene reduced average molecular weight of Polymer Compound 13 was Mn=9.0x10<sup>4</sup>, and Mw=2.0x10<sup>5</sup>.

Example 17

45 <Synthesis of Polymer Compound 14>

[9240] According to the procedure of Example 6, it was synthesized using Compound E 300mg, Compound G 64mg, N. N-bis (4-brompheny)-N-N-bis (4-brompheny)-N-Dis (4-brom

[0241] This copolymer is referred to as Polymer Compound 14. The polystyrene reduced average molecular weight of Polymer Compound 14 was Mn=3.2x10<sup>4</sup>, and Mw=6.3x10<sup>4</sup>.

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Synthetic Example 18 (Synthesis of Compound H)

[0242]

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HO OH

Compound H

[0243] Under an inert atmosphere, Compound E 1.0g was added to dichloromethane 11ml, and after completely dissolved in it, 3.5ml of boron tribromide (1.0 mol/l dichloromethane solution) was added dropwise.

[0244] After the dropwise addition, it was heated to 40°C. and stirred for 5 hours. The reaction mixture was poured into 0°C ion-exchanged water, and stirred for 1 hour. The deposited solid was filtrated and dried in vacuo, 0.80g of the desired product was obtained. (96% of vield, 99% of purity).

<sup>1</sup>H-NMR (300MHz / (CD<sub>3</sub>)<sub>2</sub>SO) : δ 7.60 (s, 2H), 8.16 (s, 2H), 10.485 (br, 2H) MS(APCI(-)):(M-H)- 373

W3(AFCI(-)).(W-H)- 373

Example 19 (Synthesis of Compound J)

[0245]

Compound J

Compound H 5.00g, 1-bromo(4-pentylphenoxy propane) 11.4g, potassium carbonate 9.24g and DMF 78ml were charged into a reaction vessel, and refuxed with heating for 4.5 hours. It was separated with bluene and water, and the organic layer was extracted, and dried with sodium sulfate. After removing the solvent, filtrated through a silica gel short column with bluene, washing is carried out twice with hot hexane, and 7.77g of the desired products was obtained. The yield was 74%, and the purity was 100%.

δ 0.88 (t, 6H), 1.24-2.55 (m, 20H), 4.25 (t, 4H), 4.34 (t, 4H), 6.86 (dd, 4H), 7.07 (dd, 4H), 7.49 (s, 2H), 7.94 (s, 2H)

Example 20

<Synthesis of Polymer Compound 15>

[0246] Compound J 391mg and 2,2-bipyridyl 130mg were dissolved in 14mL of dehydrated tetrahydrofuran, and under nifrogen atmosphere, bist [5-cyclocotadienophicke] (0 [NNI(COD)<sub>2</sub>) 250 mg was added to hits solution, and the temperature was raised to 50°C, and reacted for 3 hours. After the resction, this reaction mixture was cooled to a room temperature, added droywise to a mixed solution of 25% aqueous ammonia 7ml/methanol 50ml / 40ml ion-exchanged water, and stirred for 1 hour. The deposited prespitate was filtrated, dired under reduced pressure for 2 hours, and dissolved in tolune 20 ml. 1N hydrochlorio-acid 20mL was added and stirred for 3 hours, then the aqueous layer was removed and 4% aqueous ammonia 20mL was added to the organic layer, and after stirring for 3 hours, the aqueous layer was removed. The organic layer was added dropwise to methanol 100mL, and stirred for 1 hour, the deposited precipitate was filtrated and dried under reduced pressure for 2 hours, and then dissolved in bloune 20ml. Then,

purification through alumina column (10g of alumina) was carried out, and recovered toluene solution was added to methanol 100ml, stirred for 1 hour, and deposited precipitate was filtrated and dried under reduced pressure for two hours. A polymer was obtained in yield of 220mg. This polymer is referred to as Polymer Compound 15.

[0247] The polystyrene reduced average molecular weight of Polymer Compound 15 was Mn=2.3x10<sup>5</sup>, and Mw=1.1x10<sup>6</sup>.

Example 21

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<Synthesis of Polymer Compound 16>

[0248] It was synthesized according to the procedure of Example 20, using compound J 274mg, and N,N-bis (4-bromopheny)-N,N'bis(4-butylpheny)-1,4-phenylenedlamine 102mg. The yield of the resultant copolymer was 50mg, in the copolymer, the ratio of the repeating units of dibenzothiophene derivative and the amine derivative is 70:30. (2049) This copolymer is referred to as Polymer Compound 16. The polystyrene reduced average molecular weight of Polymer Compound 16 was Mme-1,101/4, and Mws-5,610/6.

Example 22

<Fluorescence characteristics>

on

[0250] 0.24/% chloroform solutions of Polymer Compounds 1 to 16 were spin-coated on quartz, and thin fillmuroren were prepared, respectively. The diverseence spected to the thin fillins were measured using a spectopholorometer (Hitach 150). All of them had strong fluorescence and showed fluorescence peak wavelength and fluorescence intensity as abovan 1 Table 1, respectively.

Table 1

| Polymer Compound | Fluorescence peak (nm) | Fluorescence intensity |
|------------------|------------------------|------------------------|
| 1                | 426                    | 0.67                   |
| 2                | 474                    | 1.65                   |
| 3                | 470                    | 1.62                   |
| 4                | 450                    | 4.13                   |
| 5                | 440                    | 1.82                   |
| 6                | 458                    | 3.96                   |
| 7                | 468                    | 2.89                   |
| 8                | 461                    | 2.14                   |
| 9                | 472                    | 1.73                   |
| 10               | 438                    | 1.76                   |
| 11               | 480                    | 2.18                   |
| 12               | 474                    | 3.32                   |
| 13               | 418                    | 1.38                   |
| 14               | 476                    | 2.92                   |
| 15               | 403                    | 1.94                   |
| 16               | 469                    | 3.37                   |

Example 23

<Device characteristics>

[0251] On a glass substrate on which ITO film was formed in a thickness of 150nm by sputtering method, a film was formed by a thickness of 50nm with a spin coat using a solution (Bayer Co., Baytron P) of poly (ethylenedioxythiophene)

/ polystyrene sulfonic acid, and then it was dried at 200°C for 10 minutes on a hot plate. Next, films were formed by spin-coating with a rotational speed of 800 pms, using toluene solutions of the polymer compounds obtained above whose concentration were set to being 1.5 M/S. The film thickness was about 60 ms. Purthermore, after drying this at 80°C under reduced pressure for 1 hour, an EL dvice was fabricated, by depositing about 4mm of LiF as the cathode buffer layer, about 5mm of calcium, and subsequently, about 80 mm of aluminum as the cathode. Here, after the vacuum degree reached to 1x10°P are of ses, the deposition of metal were started.

[0252] By applying a voltage to the resultant devices, EL and the maximum EL efficiency were obtained from these devices as shown in the below Table 2. The intensity of EL light emission was approximately proportional to the current density.

| Table 2          |              |                                  |  |
|------------------|--------------|----------------------------------|--|
| Polymer Compound | EL Peak (nm) | Light Emitting Efficiency (cd/A) |  |
| 2                | 480          | 0.12                             |  |
| 4                | 452          | 0.55                             |  |
| 5                | 428          | 0.02                             |  |
| 6                | 472          | 0.03                             |  |
| 7                | 460          | 0.07                             |  |
| 9                | 460          | 0.36                             |  |
| 10               | 412          | 0.31                             |  |
| 11               | 492          | 0.37                             |  |
| 12               | 472          | 0.60                             |  |
| 13               | 436          | 0.37                             |  |
| 14               | 468          | 0.28                             |  |
| 15               | 428          | 0.50                             |  |
| 16               | 480          | 1.85                             |  |

Synthetic Example 6 (Synthesis of Compound K)

35 [0253]

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## Compound K

[0254] Under an inert atmosphere, benzofuren (23.2g, 137.9mmol) and acetic acid (232g) were charged into a 1L of three-necked flask. After dissolving by stirring at a room temperature, the temperature was raised to 75°C. [0255] After the temperature rising, diluted bromine (92.6g, 579.3mmol) with acetic acid (54,9) was added dropwise.

[U295] After the temperature issuing, culticated promine (WZ.5g, 5/43-mmol) with acetic acid (E4g) was acided origined. After the dropwise addition, it was stirred for 3 house at the fixed temperature, and then left to cool. After checking the disappearance of raw material by TLC, aqueous sodium thiosulfate solution was added to terminate the reaction, and then, it was stirred at a room temperature for 1 hour. After the stirring, the cake was filterated, and further vershed with aqueous sodium thiosulfate, and water, and then dried. The resultant crude product was recrystallized in hoxane, and the desired product was obtained. (Amount 21.8g, Vield 49%)

<sup>1</sup>H-NMR (300MHz / CDCl<sub>4</sub>): δ 7.44 (d, 2H), 7.57 (d, 2H), 8.03 (s, 2H)

Synthetic Example 7 (Synthesis of Compound L)

[0256]

10

HO

## Compound L

[0257] Under an inert atmosphere, Compound K (16.8g, 50.9mmol) and tetrahydrofuran (293g) were charged into a 500mi four-necked flask, and it was cooled to -78°C. After adding dropwise n-butyllithium (80ml <1.5 mol/L hexane solution-127.3mmol), it was stirred for 1 hour at the fixed temporature.

[0258] Under an inert atmosphere, trimethoxyboronic acid (31.7g, 305.5mmol) and tetrahydrofuran (250ml) were put into a 1000ml four-necked flask, and cooled to -78°C, the above-mentioned reaction mixture was added dropwise thereon.

[0259] After the dropwise addition, the temperature raised slowly to a room temperature, and it was strred at a room temperature for 2 hours, and the disappearance of the raw material was checked by TL. Che neaction mixture was charged into concentrated sulfuric acid (309) and water (600ml) in a 2000ml beaker, and the reaction was terminated. Tolume (500ml) was added, and the organic layer was extracted, and further washed with adding water. After distilling off the solvent, 8g of thereof and ethyl acetate (160ml) were put into a 300ml four-necked flask, then, 30% autocus hydrogen persons (of 7.09g) was added, it was stimed at 40°C for 2 hours. This reaction mixture was washed with an aquous solution of an ammonium inor(fl) sulfate (7/1g) and water (500ml) in a 1000ml beaker. After stirring, the organic layer was softracted, and washed with water.

By distilling off the solvent, a crude product 7.57g of Compound L was obtained.

MS spectrum : [M-H]- 199.0

Synthetic Example 8 (Synthesis of Compound M)

[0260]

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## Compound M

[0261] In a 200ml four-necked flask, under an inert atmosphere, Compound L (2.28 g, 11.4mmol) which was synthesized according to the procedure of Synthetic Example 7 and N N-dimethylformamide (23g) were put in, after dissolving by selfring at a room temperature, potassium carbonate (0.45g, 68.8mmol) was added, and the temperature was raised to 100°C. After the temperature raising, n-octylformide (6.6g, 3.42mmol) diluted with N-N-dimethylformadide (11g was added drowbee. After the dropwise addition, the temperature was raised to 60°C, it was stirred for 2 hours at the fixed temperature, and the disappearance of the raw material was checked by TLC. Water (50ml) was added to terminate the reaction, and then toluren (50ml) was added the organic layer was extracted, and then washed with water three. After being dried with anhydrous sodium sulfate, the solvent was distillated off. By purifying the resultant crude product with a silica gel column, the desired product was obtained. (Amount: 1.84g, Yield:38%)

Example 24 (Synthesis of Compound N)

[0262]

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## Compound N

[0283] In a 500mf [our-necked flask, under an inert atmosphere, Compound M (7.50, 1.7.7mm0) which was syn-heasted according to the procedure of Synthetic Example 3 and N.4. which theyltomramide were put in, after dissolving by stirring at a room temperature, it was cooled with an ice bath. After cooling, N-bromosuccinimide (6.38g, 35.9mmo) diluted with NIA-dimethyl/comamide (225m) was added dropwise. After the dropwise addition, the temperature was explicated in the interest of the cooling of the

#### 25 Example 25

<Synthesis of Polymer Compound 17>

[0264] It was synthesized according to the procedure of Example 20, using Compound N 306mg, N N-bia (4-bromophemy)-N,N-bia(4-n-butybhemy) -1,4-phemylenediamine 153mg. The yield of the resultant polymer was 110mg. In the copolymer, the ratio of the repeating units of the dibenzoluran derivative and the amine derivative is 70:30. (0265) This polymer is referred to as Polymer Compound 17. The polystyrene reduced average molecular weight of Polymer Compound 17 was Mhm-3xt0<sup>4</sup>, and Mhw-3xt0<sup>4</sup>.

#### 35 Example 26

<Fluorescence characteristics>

[0266] According to the procedure of Example 22, a thin-film of polymer compound 17 was formed and the fluorescent light was measured. A strong fluorescence was shown and fluorescence peak wavelength was shown at 474nm.

#### Example 27

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<Device characteristics>

[0287] An EL device was fabricated according to the procedure of Example 23 using Polymor Compound 17, By applying a voltage to the resultant device, EL from this device having a peak at 488m was observed. The intensity of EL luminescence was approximately proportional to the current density. As for the device, the luminance exceeded 1 cdm<sup>2</sup> at about 3.3 V and the maximum efficiency of 1.01 cdd/was shown.

## Example 28

<Synthesis of Polymer Compound 18>

5 [0268] It was synthesized according to the procedure of Example 20, using Compound N 285mg, N, N-bis (4-bromophemy)-N, N-bis (4-b-buty) phemy)-1,4-beougher damine 48mg, and N,N-bis (4-bromophemy)-4-sisobuty-1 anilline 64mg. The yield of the resultant polymer was 90mg. In the copolymer, the ratio of the repealing units of the diberazoluran derivative and the amine derivatives is, respectively, 7b:10:20 in the order of the bromo compounds

described as the above.

[0269] This polymer is referred to as Polymer Compound 18. The polystyrene reduced average molecular weight of Polymer Compound 18 was Mn=2.8x10<sup>4</sup>, and Mw=1.7x10<sup>5</sup>.

#### 5 Example 29

< Fluorescence characteristics >

[0270] A thin film of Polymer Compound 18 was formed according to the procedure of Example 22. The fluorescence spectrum of this thin film was measured using a spectrophotofluorometer (Fluorolog made by JOBINYVON-SPEX Co.). A strong fluorescence peak was shown at 45mm.

## Example 30

#### 5 < Device characteristics>

[0271] An EL device was fabricated eccording to the procedure of Example 23 using Polymer Compound 18, By applying a voltage to the resultant device, EL luminescence from this device having a peak at 486m was observed. The Intensity of EL luminescence was approximately proportional to the current density. As for the device, the luminance exceeded 1 cuff<sup>2</sup> at about 33 V, and the maximum efficiency of 29 God Way as shown.

## Example 31

<Synthesis of Polymer Compound 19>

[0272] It was synthesized according to the procedure of Example 20, using Compound N 263mg. The yield of the resultant polymer was 50mg.

[0273] This polymer is referred to as Polymer Compound 19. The polystyrene reduced average molecular weight of Polymer Compound 19 was Mn=8.9x10<sup>4</sup>, and Mw=1.9x10<sup>5</sup>.

### Example 32

<Fluorescence characterístics >

5 [0274] A thin film of Polymer Compound 19 was fabricated according to the procedure of Example 22 and the fluorescent light was measured according to the procedure of Example 29. A strong fluorescence was observed and shows fluorescence peak at 411m.

### Example 33

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<Device characteristics>

[0275] An EL device was fabricated according to the procedure of Exemple 23 using Polymer Compound 19, 99, applying a voltage to the resultant device, EL hardya paek at 456m was observed. The intensity of EL was approxlimately proportional to the current density. As for the device, the luminance exceeded 1 cd/m² at about 4.9 V, and the maximum efficiency of 0.84 cd/4 was shown.

Synthetic Example 9 (Synthesis of Compound P)

[0276]

10

MeO MeO OMe OMe

Compound P

[0277] A starting material of 2, 2',5,5'-tetramethoxy-1, 1'-biphenyl was synthesized from 1-bromo-2, 5-dimethoxy-benzene by a coupling reaction using zero-valent nickel.

[0278] Under an lend atmosphere, 2.2.6.5/tetramethoxyr.1/bbhenyl (7.0g, 26mmmol) was charged into a threenecked fask, and dissolved in dehydrated N.N-dimethylformamide (100ml). While cooling the flask by an ice bath, from a dropping funnel, N-chilorosuccinimide (6.8g, 52mmol) of dehydrated N.N-dimethylformamide (70ml) solution was added dropwise for 15 minutes. After the dropwise addition, the temperature raised slowly to a room temperature with stirring slowly, and it was stirred for 1 day. Water (300ml) was added to the reaction mixture, and the deposited precipitate was collected. The resultant procipitate was recrystallized from toluene/hexane, and the desired product was obtained. (Amount 5.80)

<sup>1</sup>H-NMR (300MHz / CDCl<sub>3</sub>): δ 3.74 (s, 6H), 3.87 (s, 6H), 6.85 (s, 2H), 7.02 (s, 2H)

Synthetic Example 10 (Synthesis of Compound Q)

[0279]

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HO HO CI

## Compound Q

40 [0280] Under an inert atmosphere, Compound P (5.8g, 17mmmol) was charged into a three necked flask, and dissolved in dehydrated methylene chloride (100ml). While cooling the flask by an ice bath, methylene chloride solution (1 mol/L, 50ml) of boron thromdie was added droowise for 90 minutes, from a drooping funner.

[0281] Being extracted with ethyl acetate from the reaction liquid, the organic layer was washed with water, then the solvent was distilled off and the desired product was obtained. The resultant precipitate was recrystallized from toluene/ hexane, and the desired product was obtained. Amount 4.9.

1H-NMR(300MHz / CDCI<sub>0</sub>); δ 6.64 (s, 2H), 6.82 (s, 2H), 8.9 to 9.1 (br, 2H) and 9.37 (s, 2H)

Synthetic Example 11 (Synthesis of Compound R)

[0282]

10

HOO

## Compound R

15 (2833) Under an inert atmosphere, Compound Q (4.8g, 17mmmol), zeolite (6.7g, Zeolite HSZ 380+UA (Toschi)) and od-chichoroborance (170m) dived with molecular selves were charged into a three necked flask. It was stirred for 13 hours, with heating by an oil bath (180°C of the bath temperature). The reaction mixture is cooled to about a room temperature, and hexane, Goolm) was added to it. The deposited precipitate was fitted usabed with hexane, and office. The precipitate was extracted with ethylacotate, and the solution was fittrated through a short column of silica oal. The solvent was extracted with ethylacotate, and the solution was fittrated through a short column of silica oal. The solvent was destilled off, and the desired croudch was obtained.

MS spectrum : [M-H]- 267.0

(Amount 3.5a).

Synthetic Example 12 (Synthesis of Compound S)

[0284]

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F<sub>3</sub>Q 0 0 5

## Compound S

[0285] Under an inert almosphere, Compound R (4.2g), and 4-N.N-dimethylaminopyridine (5.7g) were charged into a three necked flask, and dissolved in dehydrated methylene chloride (40ml). With cooling the flask by an ice barritinoromethene sulforine acid anylydride (11g) was added dropwise for 30 minutes. Then, the temperature was raised slowly, to a room temperature, it was stirred for 5 hours. Toluene was added and filtrated by filter paper, the filtrate was filtrated through a silice gel short column, and the solvent laws a distilled off. The resultant solid was recrystallized from toluene? In laws, and the desired product was obtained. (7.8g.)

1H-NMR(300MHz / CDCl<sub>3</sub>): δ 7.80 (s, 2H) and 7.93 (s, 2H)

Example 34 (Synthesis of Compound T)

[0286]

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Compound T

[0287] Under an inert atmosphere, Compound S (3.9g), 4-n-buty/benzene boric acid (3.9g), potassium carbonate (5.1g), and tetrakis(triphenry)phosphine) palladium (0.14g) were charged into 100ml reaction tube, and dissolved in degassed tolune (20ml) and twere (20ml) and water (20ml), and twen reacted at 100°C for 12 hours. After the reaction, the tolune layer was washed with water, and filtrated thorough a silica gel short column, and the solvent was distilled off.

[0288] The resultant solid was purified through a silica gel column, and the desired product was obtained. (3.3g) 1H-NMR (300MHz / CDCI<sub>b</sub>):

δ 0.97 (t, 6H), 1.42 (m, 4H), 1.67 (m, 4H), 2.69 (t, 4H), 7.28 (d, 4H), 7.39 (d, 4H), 7.70 (s, 2H), and 7.84 (s, 2H).

25 Example 35

<Synthesis of Polymer Compound 20>

[0289] It was synthesized according to the procedure of Example 20, using Compound T 297mg, N,N'-bis(4-bromo phenyl)-N,N'-bis(4-n-butylphenyl)-1,4-phenylenediamine 138mg.

[0290] The yield of the resultant polymer was 234mg. In the copolymer, the ratio of the repeating units of the dibenzofuran derivative and the amine derivative is 70:30.

[0291] This polymer is referred to as Polymer Compound 20. The polystyrene reduced average molecular weight of Polymer Compound 20 was Mn=4.3x103, and Mw=8.2x103.

Synthetic Example 13 (Synthesis of Compound U)

[0292]

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#### Compound U

[0283] 2.2:,5-1etramethoxy-1,1-bjbhenji was brominated according to the procedure of Synthetic Example 9 with using N-bromosuccimide instead of N-birosoucciminide, and demethylation was performed according to the procedure of Synthetic Example 10, to obtain the desired product. H-NINR [000MHz] c DCDL;

δ 6.74 (s, 2H), 6.97 (s, 2H), 8.97 (s, 2H), 9.45 (s, 2H).

Synthetic Example 14 (Synthesis of Compound V)

[0294]

HO O

Compound V

[0295] According to the procedure of Synthetic Example 11, Compound U was used instead of Compound Q, and the desired product was obtained.

MS spectrum: (M-H1 356, 9.

Example 36 (Synthesis of Compound W)

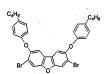
20 [0296]

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Compound W

[0237] In a three necked flask, Compound V (0.83q), 4-but/pheny/boronic soid (1.6g), and copper(II) seatate (0.71g) molecular slews 4.0 (9.3, powder) and dielohromathen (31ml) and pyridine (0.41g) were stirred at a room temperature for 3 hours in air. The reaction mixture was filtrated through certie, and chloroform and hydrochloric soid (1 mol). I were added to the filtrate, the organic layer was extracted, and dried with sodium suitate. It was purified frough a slice age lookum (foluene:cyclohoxane=12), washed with hot haxane, then the desired product (white solid) was obtained, (0.82a).

<sup>1</sup>H-NMR(300MHz / CDCl<sub>3</sub>):

δ 0.92 (t, 6H), 1.35 (m, 4H), 1.54 (m, 4H), 2.58 (t, 4H), 6.89 (dd, 4H), 7.12 (dd, 4H), 7.37 (s, 2H), and 7.84 (s, 2H)

45 Example 37

<Synthesis of Polymer Compound 21>

[0289] It was synthesized according to the procedure of Example 20, using Compound W 174mg, N.N-bis (4-homopheny)-N. bis (4-ho-hybpeny)-1.4-phenylenediamine 28-pg... The yield of the resultant polymer was 95mg. In the oppolymen, the ratio of the repeating units of the dibenzofuran derivative and the armine derivative is 70:30 in the above described order of the corresponding brone compounds. This polymer is referred to as Polymer Compound 21. [0299] The polystyrene reduced average molecular weight of Polymer Compound 21 was Mn-1.6x10<sup>4</sup>, and Mw-4.7x10<sup>4</sup>. Example 38 (Synthesis of Compound X)

T03001

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Compound X

[3301] In a three neckod flask under an inert amnosphere, Compound V (2.5g) was dissolved in N.N-dmethylformaindie (54m). To this, 3-bromopropylbenzee (4.2g) and potassium contonate (4.3g) were added, and then, the temperature was raised to 160°C, and siltred for 6 hours. Then, tokene was added to 1s, and washed with dilute hydrochloric acid and water. The organic layer was dried with southin sulfato, and the solvent was desilled off. The resultant solid was purified by silica gel column chromatography using hexane/ chloroform, and the desired product was obtained. (3.8a).

<sup>1</sup>H-NMR(300MHz / CDCl<sub>3</sub>):

δ 2.18-2.27 (m, 4H), 2.93 (t, 4H) and 4.10 (t, 4H), 7.17 to 7.32 (m, 12H) and 7.70 (s, 2H)

Example 39

<Synthesis of Polymer Compound 22>

[0902] It was synthesized according to the procedure of Example 20, using Compound X 291mg, N.N-bid.

(4-bromophemy).N.N-bid(4-huy)theny)-1,4-phoryindealmen is 13mg. The yield of the resultant polymer was
120mg. In the copolymer, the ratio of the repeating units of the dibenzofuran derivative and the amine derivative is 70:
30 in the above described order of the corresponding bromo compounds.

[0303] This polymer is referred to as Polymer Compound 22. The polystyrene reduced average molecular weight of Polymer Compound 22 was Mn=3.5x10<sup>4</sup>, and Mw=1.4x10<sup>5</sup>.

Example 40

<Pre><Pre>conduction of Polymer Compound 23>

[0304] 2mg of Polymer Compound 13, and 0.1 mg of Ir complex, Blp\_lr(acac) were dissolved in 0.2ml of toluene, and a film was formed in a thickness of about 200nm by spin coating, and dried at a room temperature in a reduced pressure, overnight.

[0305] The fluorescence spectrum of the resultant thin film was measured, and phosphorescence emission from Ir complex was observed.

[0306] The polymer compound of the present invention having thiophene-condensed ring unit a new polymer compound which can be used as a light-emitting material, a charge transporting material, etc.

Claims

 A polymer compound having a polystyrene reduced number average molecular weight of 10<sup>3</sup>-10<sup>8</sup>, and comprising a repeating unit of formula (1-1) or (1-2):

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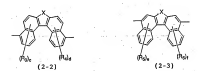
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wherein, Ring A, Ring B, Ring C, and Ring D, each independently represent an aromatic ring: X represents S or O; and these repeating units may have a substituent selected from the group consisting of an alkyl group, alkoyd group, alkything group, anyl group, anyloxy group, anything group, anylalkyl group, anylalkoyn group, anylalkynting group, anylalkenyl group, anylalkynyl group, anino group, substituted amino group, silyl group, substituted silyl group, halpena dom, anyl group, anyloxy group, and mino group.

- A polymer compound according to claim 1, wherein Ring A, Ring B, Ring C, and Ring D are aromatic hydrocarbon rings.
  - 3. A polymer compound according to claim 1 or 2, wherein Ring A and/or Ring B, and Ring C and/or Ring D have at least one substitutent selected from an allyl group, allowg group, allyling group, anyling group, and group, anyling group, and group, anyling group, and group, anyling group, and group.
- A polymer compound according to claim 1, wherein the repeating unit of formula (1-1) is a repeating unit selected from formulae (2-1) to (2-5):

$$(R_1)_a$$
  $(R_2)_b$ 





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wherein, X represents S or C; R<sub>1</sub>, R<sub>2</sub>, B<sub>3</sub>, R<sub>4</sub>, R<sub>5</sub>, R<sub>5</sub>, R<sub>6</sub>, R<sub>6</sub>, and R<sub>10</sub>, each independently represent an ality group, alktying orgue, alfything orgue, and group, and group

A polymer compound according to any one of claims 1 to 4, wherein said polymer compound further comprises a repeating unit of formula (3), formula (4), formula (5), or formula (6):

wherein,  $A_1$ ,  $A_2$ ,  $A_3$ , and  $A_4$ , each independently represent an aylene group, a division the terrocyclic group, or a division throughout structure;  $X_1$ ,  $X_2$ , and  $X_3$  each independently represent  $CR_{11} = CR_{12}$ , C=C-1,  $V(R_{12})$ , or  $\{SR_{11}R_{12}\}$ ;  $R_{11}$  and  $R_{12}$  each independently represent a hydrogen atom, alkyl group, arry group, monovalent heterocyclic group, carboxyl group, substituted carboxyl group, or group group,  $R_{13}$ ,  $R_{14}$ , and  $R_{12}$  each independently represent a hydrogen atom, alkyl group, any group, monovalent heterocyclic group, any lalkyl group, and substituted amino group; if represents an integer of  $C_2$ ; it represents an integer of 1-12; when  $R_{11}$ ,  $R_{12}$ ,  $R_{12}$ ,  $R_{13}$ , and  $R_{14}$  exist in plural. Respectively, a plurality of them may be the same or different.

 A polymer compound according to claim 5, wherein the repeating unit of formula (3) is a repeating unit of formula (7), (8), (9), (10), (11), or (12):

wherein, R<sub>16</sub> represents an alkyl group, alkoxy group, alkylthio group, anyl group, anylton group, anylthio group, anylakyl group, anylakoxy group, anylakylthio group, anylakenyl group, anylakynyl group, amine group, sichidad amine group, silvl group, substituted silvl group, halogen atom, acyl group, acyloxy group, immo group, amide group, imide group, monovalent heterocyclic group, carboxyl group, substituted carboxyl group, or cyano group; I represents an integer of 0-4, when R<sub>16</sub> exists in pluria, a plurality of them may be the same or different,

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wherein, R<sub>17</sub> and R<sub>16</sub> represent each independently an aikyl group, alkoxy group, allythio group, any group, and group, any group, and group,

$$\begin{array}{c} \begin{pmatrix} R_{19} \\ P_{20} \end{pmatrix} & P_{20} \\ \vdots \\ R_{21} \\ R_{22} \end{pmatrix}_{p} & (9) \end{array}$$

wherein,  $H_{19}$  and  $H_{22}$  each independently represent an alkyl group, alkoxy group, alkylink group, any flaving represent an alkyl group, any flaving group, and group, any flaving group, and group, any flaving group, and group group group, and group group group, and group group

$$-\left(A_{15}\right)_{x} \left(A_{16}\right)_{y} \left(A_{16}\right)_{y}$$

wherein, R<sub>23</sub> represents an alkyl group, alkoxy group, alkylthio group, aryl group, aryloxy group, arylthio group, arylalkyl group, arylalkoxy group, arylalkylthio group, aryl alkenyl group, arylalkynyl group, amino group, substituted

amino group, silyl group, substituted silyl group, halogen storm, acyl group, acyloxy group, imino group, amide group, minde group, minde group, minde group, minde group, acyloxy group,

wherein, R<sub>24</sub> and R<sub>25</sub> each independently represent an alkyl group, alkoxy group, alkylthio group, aryl group, arylchio grou

wherein,  $R_{30}$  and  $R_{35}$  each independently represent an alkyl group, alkoxy group, alkylthio group, anyl group, anylong group, anylon

A polymer compound according to claim 5, wherein the repeating unit of formula (4) is a repeating unit of formula (13):

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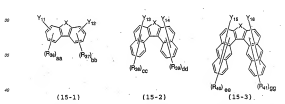
wherein,  $A_{16}$ ,  $A_{17}$ ,  $A_{18}$ , and  $A_{19}$  each independently represent an anylene group or a divalent heterocyclic group;  $A_{16}$ ,  $A_{11}$ , and  $A_{12}$  each independently represent an anyl group or a monovalent heterocyclic group;  $A_{16}$ ,  $A_{17}$ ,  $A_{18}$ ,  $A_{19}$ , and  $A_{19}$  may have a substituent; v and w each independently represent an integer of 0 or 1, and 0-e-ww-c1.

 A process for producing a polymer compound according to any one of claims 1 to. 7, wherein condensation polymerization is carried out using a compound of formula (14) as one of the raw materials:

(13)

wherein, U represents a repeating unit represented by the above formula (1-1) or (1-2);  $Y_1$  and  $Y_2$  each independently represent a substituent capable of condensation polymerization.

 A process according to claim 8, wherein the compound of formula (14) is a compound of formula (15-1), (15-2), or (15-3):



wherein, X represents S or O,  $Y_{11}$ ,  $Y_{12}$ ,  $Y_{13}$ ,  $Y_{12}$ ,  $Y_{13}$ , and  $Y_{16}$  each independently represent a substituent capable of condensation polymerations,  $P_{B_0}$ ,  $R_{B_1}$ ,  $R_{B$ 

10. A process according to claim 8 or 9, wherein condensation polymerization of a compound of one of the below formulae (16) to (19) in addition to the compound of formula (14), is carried out:

$$Y_3-Ar_1-Y_4$$
 (16)

$$Y_5 - \{Ar_2 - X_1\}ff - Ar_2 - Y_6$$
 (17)

$$Y_7 - Ar_4 - X_9 - Y_8$$
 (18)

wherein, Ar<sub>1</sub>, Ar<sub>2</sub>, Ar<sub>3</sub>, Ar<sub>4</sub>, If, X<sub>1</sub>, X<sub>2</sub>, and X<sub>3</sub> are the same as above; Y<sub>3</sub>, Y<sub>4</sub>, Y<sub>5</sub>, Y<sub>6</sub>, Y<sub>7</sub>, Y<sub>8</sub>, Y<sub>9</sub>, and Y<sub>10</sub> each independently represent a substituent capable of condensation polymerization.

- 11. A process according to claim 9 or 10, wherein Y<sub>1</sub>, Y<sub>2</sub>, Y<sub>3</sub>, Y<sub>4</sub>, Y<sub>5</sub>, Y<sub>6</sub>, Y<sub>7</sub>, Y<sub>8</sub>, Y<sub>9</sub>, and Y<sub>10</sub> each independently represent a halogen atom, alkyl sulfonate group, anyl sulfonate group, carylakilyl sulfonate group; and condensation polymerization is carried out in the presence of a nickel zero-valent complex.
- 12. A process according to any one of claims 9 to 11, wherein Y<sub>1</sub>, Y<sub>2</sub>, Y<sub>3</sub>, Y<sub>4</sub>, Y<sub>5</sub>, Y<sub>6</sub>, Y<sub>7</sub>, Y<sub>8</sub>, Y<sub>3</sub> and Y<sub>10</sub> each independently represent a halogen atom, alkyl sulfonate group, anyl sulfonate group, anylalkyl sulfonate group, boric acid group or boric ester group;

the ratio of the total number of moles of the halogen atom, alkyl sulfonate group, anyl sulfonate group, and anylalkyl sulfonate group, to the total number of moles of boric acid group and boric ester group is about 1, and condensation polymerization is carried out using nickel or a palladium catalyn.

A compound of formula (15-1), (15-2), or (15-3) as defined in claim 9.

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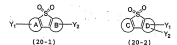
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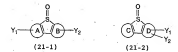
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14. A process for producing a compound of formula (14) as defined in claim 8, wherein a compound of formula (20-1) or (20-2):



wherein, Ring A, Ring B, Ring C, and Ring D, are as defined in claim 1 for formulas (1-1) and (1-2); and  $Y_1$  and  $Y_2$  are the same as those in formula (14), is reduced using a roducing agent.

 A process for producing a compound of formula (14) as defined in claim 8, wherein a compound of formula (21-1) or (21-2):



wherein, Ring A, Ring B, Ring C, and Ring D, are as defined in claim 1 for formulas (1-1) and (1-2); and  $Y_1$  and  $Y_2$  are the same as those in formula (14). It is reduced using a reducing agent.

16. A composition comprising one or more compounds selected from a hole transporting material, an electron trans-

porting material, and light-emitting material, and one or more polymer compounds according to any one of claims

- 17. An ink composition comprising at least one polymer compound according to any one of claims 1 to 7.
- An ink composition according to claim 17, which has a viscosity of 1 to 20mPa-s at 25°C.

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- 19. A light emitting thin film comprising the polymer compound according to any one of claims 1 to 7.
- A conductive thin film comprising the polymer compound according to any one of claims 1 to 7.
  - 21. An organic semiconductor thin film comprising the polymer compound according to any one of claims 1 to 7.
- 22. A polymer light emitting device comprising an organic layer between electrodes consisting of an anode and a cathode, wherein said organic layer comprises the polymer compound according to any one of claims 1 to 7.
  - 23. A polymer light emitting device according to claim 22, wherein the organic layer is a light emitting layer.
- 24. A polymer light emitting device according to claim 23, wherein the light emitting layer further comprises a hole transporting material, an electron transporting material, or a light-emitting material.
- 25. A flat light source comprising a polymer light emitting device according to any one of claims 22 to 24.
- 26. A segment display apparatus comprising a polymer light emitting device according to any one of claims 22 to 24.
- 27. A dot-matrix display apparatus comprising a polymer light emitting device according to any one of claims 22 to 24.
- 28. A liquid crystal display comprising a polymer light emitting device according to any one of claims 22 to 24 as a back light.



# PARTIAL EUROPEAN SEARCH REPORT

Application Number

which under Rule 45 of the European Patent Convention EP 03 25 1503 shall be considered, for the purposes of subsequent proceedings, as the European search report

DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document with indication, where appropriate, Relevant to claim CLASSIFICATION OF THE APPLICATION (Int.CI.7) Category of relevant passages US 5 876 864 A (KIM CHUNG YUP ET AL) 6,7, 9-13, C08G61/12 2 March 1999 (1999-03-02) H05B33/14 19-28 C09K11/06 \* examples 15-18; table 1 \* SHIMOMURA, SATO:
"S-Alkyldibenzothiophenium Salts as new
thermal latent cationic initiators" 6,7, 9-13, γ 19-28 MACROMOLECULES, vol. 31, 27 February 1998 (1998-02-27), pages 2013-2015, XP002244280 scheme 1 -& SUPPORTING INFORMATION MACROMOLECULES, pages 1-4, XP002245115 WIRTH, WAESE, KERN: "Synthese und 6,7, Eigenschaften von Oxydo-p-Oligophenylenen" 9-13, DIE MAKROMOLEKULARE CHEMIE, vol. 86, 1965, pages 139-167, XP009012341 \* page 139-167 \* 19-28 TECHNICAL FIELDS SEARCHED (Int.CL7) -/--C08G H05B C09K INCOMPLETE SEARCH The Search Division considers that the present application, or one or more of its claims, dosestion not comply with the EPO to such en extent that a meaningful search into the state of the art cannot be carried out, or can only be carried out partially, for these claims. Cleims searched completely: Claims searched incompletely: Claims not searched : Reason for the Emitation of the search. see sheet C Place of search MUNICH 16 June 2003 Marsitzky, D T: theory or principle underlying the invention
E: earlier patent document, but published on, or after the filing date
D: document cited in the application
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Y : particularly relevant if continued with another
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A : technological background
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## INCOMPLETE SEARCH SHEET C

Application Number EP 03 25 1503

Claim(s) searched incompletely: 4, 6-7, 9-14, 19-28

Claim(s) not searched: 1-3, 5, 8, 15-18

Reason for the limitation of the search:

Present claims 1-6, 8, 13-15 and 17-18 relate to an extremely large number of possible compounds and products. Support within the meaning of Article 84 EPC and/or disclosure within the meaning of Article 83 EPC is to be found, however, for only a very small proportion of the compounds and products claimed. In the present case, the claims so lack support, and the application so lacks disclosure, that a meaningful search over the whole of the claimed scope is impossible.

Consequently, the search has been carried out for those parts of the claims which spear to be supported and disclosed, mamply those parts claims which spear to be supported and disclosed, mamply those parts the mamples (p. 89-134) and in claims 4 (relating to formula 2-1 is substituted dibenzothughner-derived homopolymers, subst. dibenzothiophene-derived polymers and oxidized dibenzothiophene derived polymers, of (copolymers wherein the commonmer is selected from formula 19, 7 (copolymers relating to formula 2-1 with formula 13, 9 (process utilizing compounds of formula 15-1), 10 (process utilizing compounds of formula 15-1), and formula 16-quinox line-deriv. or formula 15-1 and formula 16-quinox line-deriv. or (relating to compound 20-1) and claims 10-28 (products derived from the previously described polymers) and copolymers.



# PARTIAL EUROPEAN SEARCH REPORT Application Number

EP 03 25 1503

|          | DOCUMENTS CONSIDERED TO BE RELEVANT                                                                                                                                                                                                                | CLASSIFICATION OF THE<br>APPLICATION (Int.CL7) |                                         |
|----------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------|-----------------------------------------|
| Category | Ottation of document with indication, where appropriate,<br>of relevant passages                                                                                                                                                                   | Relevant<br>to claim                           |                                         |
| A        | SIRRINGHAUS, FRIEND, WANG, LEUNINGER, MOLLEN: "Dibenzothienobisbenzothiophene - a novel fused ring digomer with high field-effect mobility" lower with high JONENAL OF MATERIALS CHEMISTRY, vol. 9, 1999, pages 2095-2101, XP002244281 schemes 4-5 | 4,6,7,<br>9-14,<br>19-28                       |                                         |
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# ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO. EP 03 25 1503

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. This members are as continued in the European Patent Office CDP file on The European Patent Office is in one yet label for these particulars which are merely given for this purpose of information.

16-06-2003

| :    | Patent docume<br>cited in search re | nt<br>port | Publication date |    | Patent family<br>member(s) | Publication<br>date |
|------|-------------------------------------|------------|------------------|----|----------------------------|---------------------|
| : US | 5876864                             | A          | 02-03-1999       | KR | 176336 B1                  | 01-04-1999          |
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For more details about this annex : see Official Journal of the European Palent Office, No. 12/82